THE CHEMICAL FORMULARY

THE CHEMICAL FORMULARY

A CONDENSED COLLECTION OF VALUABLE, TIMELY,
PRACTICAL FORMULAE FOR MAKING THOUSANDS
OF PRODUCTS IN ALL FIELDS OF INDUSTRY

VOLUME III

Editor-in-Chief
H. BENNETT



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H. BENNETT

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PREFACE TO VOLUME II

The gratifying reception accorded Volume I of the Chemical Formulary together with the helpful and constructive criticisms received from reviewers and chemists have manifestly proved the need for a book of this type covering modern formulation in commercial chemistry.

While Volume I is complete in itself, the Editors felt it was impossible within the scope of one book to include all the formulae compiled for the numerous subject headings in the book. Volume II therefore is not a duplication or revision of Volume I but an entirely new work giving further formulae on the subjects treated in the first volume as well as more detailed information on processes and fundamental principles involved.

It will be noticed that all patented formulae have the patent number included. A helpful article on what is patentable in chemical compounding: infringements, licensing, etc., is another important addition to the book. It must be borne in mind in this connection that patented formulae cannot be used in the manufacture of commercial products unless prior arrangements have been made with the patentee.

The Editorial Board has been considerably enlarged and consequently it has been possible to include formulae hitherto unavailable.

A certain amount of criticism was directed toward the use of trade names in Volume I. It was contended by the critics that formulae containing trade names should be eliminated regardless of their value. Considerable thought was given to this contention and it was felt that, masmuch as chemical trade-name products are being used in an ever-increasing number of formulae in every class of chemical manufacturing, these formulae should be included unless the application was exceptionally limited.

A second subject of criticism was the non-uniformity of systems of weights and measures used in the book. Since there is no uniformity in such systems in commercial practice and since the main purpose of the book is to familiarize the reader with commercial practice it was thought best not to attempt to standardize these systems.

In the Preface to Volume I, it was emphasized that the chemistry taught in schools and colleges is rightly confined to synthesis, analysis and engineering whereas in commercial manufacture many of the products so made are not synthetic or definite chemical materials but consist of mixtures, blends or highly complex compounds.

Because of the paucity or antiquity of the literature in this field and because of the difficulty encountered even by experienced chemists on entering new fields a definite need has existed for a modern compilation of formulae for chemics compounding and treatment.

In addition to an Editorial Board composed of chemists and engineers in many industries, publications, laboratories, manufacturers and individuals have been consulted to obtain the latest and best information in the numerous fields covered in the book.

It is important to remember that repeated experiments may be necessary to get the best results, especially when the field is intricate or unfamiliar. Again, although many of the formulae are being used commercially, some of them have been taken from patent specifications and the literature. Since these sources are subject to various errors and omissions, due regard must be given to this factor.

Formulae must be considered chiefly as starting points, variations have to be made to meet individual requirements and specifications. In cases of doubt or difficulty it is advisable at all times to consult other chemists or technical workers familiar with the particular field. This applies particularly in the case of the layman, as while a certain expense is involved this is more than compensated for by the saving of time, money and material.

As mentioned in Volume I it is hoped that those who have found a work of this kind helpful, will bring to our attention any errors they come across and will feel free at all times to make any constructive criticisms or suggestions.

PREFACE TO VOLUME III

Because of an insistent demand for new and additional formulae Volume III of the Chemical Formulary is being published a year in advance of original plans. In technical compounding there is no rest or "breathing spell"—no "status quo." Improvements are being made daily and new ideas and methods are continually being initiated and applied. New sources of data in many fields are being opened up in order to increase the breadth and scope of information.

As far as possible there has been included information especially requested by users of Volumes I and II. Diligent cooperation on the part of many chemists, engineers, teachers, technicians and other workers has made this possible.

The editor-in-chief wishes to thank all those who have helped in this work, which, in so short a time, has found a place as a highly useful tool and time suver at the right hand of so many technical workers. In many cases it has proved to be a veritable catalyst in stimulating new products and processes.

Any thoughts for improving succeeding volumes and any new formulae or data, will, as heretofore, be most welcome. To make reference more easy the index in this volume is inclusive of Volumes I, II and III so that three separate indices need not be consulted.

H. BENNETT

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ABBREVIATIONS

amp		·ampere	
avoir		avoirdupois	
b.p		. boiling point	
Bé		Baumé	
cc		cubic centimeter	
c.d		current density	
		chemically pure	
		free from chlorine	
		free from prussic acid	
	.		
	• • • • • • • • • • • • • • • •		
	• • • • • • • • • • • • • • • • • • • •		
L	• • • • • • • • • • • • • • • • • • • •	nter	
т.р	• • • • • • • • • • • • • • • • • • • •	meiting point	
		Hydrogen Ion Concentration	
		A quantity sufficient to mal	62
		revolutions per minute	
8p. ()		.specific gravity	
8q. dm		square decimeter	
U.S.P		.U. S. Pharmacopeia	
v		. voltage	
wt		.weight	
		=	

ADHESIVES

White Glue

A solution consisting of:	
Animal Glue	100 oz.
Zinc Oxide	50 oz.
Water	100 oz.

gives a glue which sets quite hard and is very strong.

Clua

Urea.	1	lb.
Casein	2	lb.
Hydrate of Lime	1/4	lb.

Black Albumen from Blood

Let slaughterhouse-blood stand in shallow dishes or pans, cut the blood jelly, sift the serum off. The residue is stirred in water to a paste, and put through a filter press. Evaporation in vacuum produces from the second filtrate the dark black albumen used for veneering and laminating.

"Salamyn-Plant" Glue

a. Potato-Starch

Water (35° C.)	105 1.
b. Caustic Soda (35°	Bé.) 15 kg.
c. Hydrochloric Acid	
Water	10 l.
d. Upholsterer's Glue	e 260 kg.
Sting for 3/ hour of	tor adding d Stir

35 kg.

Stir a for ¼ hour after adding d. with b until glassy, then add c.

Calcium Saccharate Glue

Water, Boiling	70 g.
Sugar	6 g.
Lime, Fresh Slaked	1.5 g.

Let stand, stir often, cover. After a few days pour off from bottom deposit, and soak in the solution,

Carpenter's Glue then warm to solution.

Marine Linoleum Cement

Decks to be covered with linoleum should be thoroughly cleaned, and the linoleum stuck to the deck with the following adhesive:

To make 10 gallons, first cut 4 oz. of crude (ham) rubber into small lumps and

dissolve in 4½ gallons of gasoline. It will require about two days to get the rubber into colloidal solution. When in proper condition it should string about two inches thumb and forefinger. Cut 19 lb. of guin shellac in 34 lb. of denatured (or wood) alcohol. Add 62 lb. of whiting then add the rubber solution. For best results this mixture should be ground in an iron or pebble mill,

Linoleum Glue

a. Rye or Barley Flour	50	kg.
Water, tepid	250	
b. Caustic Soda (20° Bé.)	20	kg.
c. Turpentine, Venice,	20-25	ko.

20-25 kg. Part a dispersed by stirrer is mixed with b (dissolved). The mixture is then boiled, and after cooling emulsified by adding c (while stirring add).

Painters' Glue (Cold)

· • <i>)</i>
350 1.
100 kg.
21 kg.
56 kg.
stirring for
and neutral-
til red color
pears (in a

Wall Size

sample). Stir 1/2 hour more.

Aluminum Stearate	4	oz.
Turpentine		oz.
Mineral Spirits (150-190° C.)	71	oz.
Heat the turpentine to 180°	F	. and
add the stearate slowly while stir	rine	z con-

tinuously. Add mineral spirits and stir until clear.

l'ainters' Size	
Potato-Starch (Air-Dried)	7.8 g.
Calcium Chloride	7.0 g.
Water	3.0 g
m	•

The aqueous paste, when compact, is dried and ground. The excess chloride can be extracted with aqueous alcohol, yielding a better paintable and quicker drying product.

Paperhanger's Paste

Use a cheap grade of rye or wheat flour, mix thoroughly with cold water to about the consistency of dough or a little thinner, being careful to remove all lumps. Stir in a tablespoonful of powdered alum to a quart of flour, then pour in boiling water, stirring rapidly until the flour is thoroughly cooked. Let this cool before using and thin with cold water.

Venetian Paste

a. White or Fish Glue	4	oz.
Cold Water	8	02.
b. Venice Turpentine	2	fl. oz.
c. Rye Flour	1	lb.
Cold Water	16	fl. oz.
d Buling Water	64	fl. oz.

Soak the 4 oz. of glue in the cold water for 4 hours. Dissolve on a waterbath (glue-pot) and while hot stir in the Venice turpentine. Make up c into a batter free from lumps and pour into d. Stir briskly, and finally add the glue solution. This makes a very strong paste, and it will adhere to a painted surface, owing to the Venice turpentine in its composition.

Flour Paste

a. Wheat Flour	2	lb.
Cold Water (1 quart)	32 fl.	oz.
b. Alum	1	oz.
Hot Water	4 fl.	oz.
c. Boiling Water	96 fl.	θZ.

Work the wheat flour into a batter free from lumps with the cold water. Dis-solve the alum as designated in b. Now stir in a and c and, if necessary, continue boiling until the paste thickens into a semi-transparent mucilage, after which stir in the solution b. This makes a very fine paste for wallpaper.

Sinclair's Glue

Formula No. 1

"Very Good" Glue or Gelatin 50 oz. Water 100 oz. Glycerin 4 or 6 oz. Thymol or Menthol 0.15 oz.

The smaller amount of glycerin is for summer or tropical use, and the larger amount for winter. Gelatin is preferable, for commercial glue varies in quality and generally requires neutralizing to litmus with weak alkali. The following is a simple test for a "very good" give. "On soaking glue in excess of cold \.`\$

water overnight, a gelatinous coherent mass is obtained, weighing, when drained, at least four times the weight of the original glue." With the very best glue a mass weighing five times the original weight is obtained.

No. 2				
Isinglass				oz.
Gelatin				υz.
Water			200	υz.
Tannic Acid			12	oz.
Glycerin	8	or	more	oz.
Menthal or Thymal			0.15	OZ.

Menthol or Thymol

Rubber

Shellac

This forms a stronger adhesive, is perhaps more elastic, and has the advantage of somewhat hardening the skin so that the tendency to blistering is almost completely eliminated.

Marine Glue 100 g. 600 g. Turpentine Coal Tar Oil 600 g. 300 g.

Warm together and mix till smooth.

Preserving Glue

Add 3 ounces of ordinary borax to each gallon of glue or add 1 ounce of formaldchyde to the gallon or 1 ounce of carbolic acid. Adding 1/2 ounce of 28% acetic acid to 2 pounds of glue will also prevent the souring and also has a tendency to make it waterproof.

Casein Glue

	Formula	No. 1			
Casein				100	oz.
Water		220	to	230	oz.
Hydrated	Lime	20	to	30	
Water				100	oz.

Water Silicate of Soda Copper Chloride 70 oz. 2 to 3 oz. 30 to 50 oz. Water

The 220 to 230 parts of water added to the casein is approximately the right amount to use with Argentine (naturally soured) casein; but if a different casein is used the water requirement will lie somewhere between 150 and 250 parts by weight. The correct amount for different caseins must be determined by trial.

The formula presupposes that a high calcium lime will be used. A lime of lower grade may be used, but a proportionately larger amount of it will be needed, or the water resistance of the glue will be sacrificed. It is suggested that for the first trial the user try 25 parts of lime. If this does not give proper results the amount can be varied within the limits specified.

The density of the silicate of soda used should be about 40° Baumé, with a silica soda ratio of from 3 to 3.25.

Copper sulphate can be substituted for

copper chloride.
Place the casein and water in the bowl of a mixing machine and rotate the paddle slowly, stirring the mixture until all the water has been absorbed and all the casein moistened. If the casein is allowed to soak beforehand it is more readily dissolved in the mixing process. Mix the hydrated lime with water in a separate container. Stir this mixture vigorously at first, but just before it is added to the casein stir just enough with a gentle rotary motion to keep the lime in suspension. Pour the milk of lime quickly into the casein.

When casein and lime are first combined they form large, slimy lumps, which are balls of dry casein coated with partly dissolved casein. These break up rapidly, becoming smaller and smaller, and finally disappear. The solution, in the meantime, is becoming thin and fluid. At this point stop the paddle and scrape the sides and bottom of the container, and then stir again. If a deposit of casein remains unacted on, it may cause

more lumps later.

When about two minutes have elapsed since the lime and casein were united, it may be noticed that the glue has begun to thicken a little. Add the sodium silicate now, or else the glue will become too thick. The glue will momentarily become even thicker, but this thickness will soon change to a smooth and fluid consistency.

Continue the stirring until the glue is free from lumps. This should not take more than 15 or 20 minutes from the time the lime was added. If the glue is a little too thick, add a small amount of water. If the glue is too thin, it will be necessary to start over again, using a

smaller proportion of water.

The copper salt may be added at any one of several times during the mixing process. If added as a powder before the casein is soaked, it may have a corrosive action upon the metal container. The copper salt, if added as a powder, should be thoroughly mixed with the casein before the addition of the lime. Copper salt may be placed in solution and conveniently stirred into the moistened casein immediately before the lime is added or after all the other ingredients have been combined. If the copper solution is added at the end of the

mixing period, pour it into the glue in a thin stream and stir the mixture vigorously. Continue stirring until any lumps, which may have formed by the coagulation of the glue and the copper smooth violet-colored glue is obtained.

Glue prepared by this formula has proved to be exceptionally strong and durable, even under wet or damp conditions.

Formula No. 2

The mixing is the same as for above formula except for the omission of the copper chloride. The glue made by this formula has a medium consistency, excellent working properties, a good working life, and makes joints of high strength, but it falls somewhat short of the previous formula in water-resisting properties, especially when the lower amounts of lime are used.

Casein	100	oz.
Water	200	oz.
Sodium Hydroxide (Caustic		
Soda)	10	
Water	50	OZ,

Bring the casein and water together according to the directions for mixing glue prepared by previous formula. Dissolve the caustic soda in water in a separate container, and while the mixing paddle is revolving sprinkle the caustic sods solution into the damp casein. Stir slowly until a thin, smooth glue has been obtained. The consistency of the finished product may be altered by adding more casein if it is too thin, or by adding water if it is too thick. Silicate of soda is sometimes added to thicken or to reduce the cost of the glue per unit of volume

This glue has exceptional strength when dry, but when exposed to moisture it weakens as rapidly as animal or vegetable glue.

Cold Glue (Cascin)

Formula No. 1

a. Casein, Dry		70	ø.
Trisodium P	hosphate	10	
Lime Hydra		20	
Sodium Fluo			g.
b. Water		200	g.
Pine Oil			g.
a is soaked wit	h <i>b</i> .		•

No. 2

a.	Casein	60 g.
	Lime, Hydrated	15 g.
	Trisodium Phosphate	4 g.

Sodium Flu	oride	4 g.	Water	50 cc.
Nut Meal	101140	17 g.	Glue Jelly	
b. Water		200 g.		- ⁵ g.
Stir a with b	: paste reads		Modern Casein Adhesi	•
minutes.	, paste ready	arter bo	Modern Casein Adhesi	ve Powders
	No. 3		For use stir with 140 tin	nes the amount
Casein		0-30 g.	of water (cold). After 1/2	to 1/4 hour, a
Caustic Soda	2	0-30 g.	homogeneous, viscous solu	tion is gotten
(36° Bé.)	0.2	-0.6 g.	ready for use.	
(00 100)		.7-2 g.	Formula No.	1
	0. 0	5.	Lactic Acid-Casein	=0
Water	79.8	-68 cc.	Marble-Lime Hydrate	70 g. 13 g.
7	No. 4		Trisodium Phosphate	5 g.
Casein		–30 g.	Sodium Fluoride	4 g.
Soda Ash		4.5 g.	Sodium Sulphate, Pure,	- 6.
Water		35.5 ee,	Anhydrous	6 g.
	No. 5		Naphtha, Refined	2 g.
Casein		-30 g.	No. 2	J
Borax	2	- 5 g.	Lactic Acid-Casein	60 g.
Water	78	-65 ec.	Slaked Lime	60 g. 20 g.
ľ	No. 6		Trisodium Phosphate	10 g.
Casein		-30 g.	Aniline	8 g.
Ammonia (sp. g		- 24 cc.	Mineral Oil	2 g.
Water		-46 ec.	No. 3	Ü
	No. 7		Lactic Acid-Casein	50 g.
Casein	12 g.]	Slaked Lime	16 g.
Borax	1.5 g.	Knead	Trisodium Phosphate	8 g.
Ammonia (0.91)		Kneard	Sodium Sulphite	8 g.
Water	85 g.	}	Sodium-Waterglass, Dry	6 g.
N	No. 8		Copper Chloride Hardwood-Meal	2 g.
Casein 2	0 g.	1	Mineral Oil	10 g.
Water 60	g. soak		Millerat Oil	11/2 g.
Disodium Hy-	σ,	i l	Air-plane Propeller	Clue
drogen Phos-	1			Mix at 15°
phate :	3 g.	Knead	1. Diack Dioou	C., stop mix-
Water 20	0 g. dissolve	. 1	g. J	ing for two
Caustic Soda			Water 6 g.	hours
(10%)	6 g.	}	Add:	365 AP
Mix all in warm	water beth		Sinked Lillie 0.00 g.	Mix until
			Water 1 g.	thiek
	Vo. 9			
Casein	20 g.		Mordant for Handles of K	itchen-Knives
Water Borax	80 g.	,, ,	a. Potassium Bienromate	15 g.
Ammonia (0.91)		Varm for	Water	1000 cc.
Caustic Soda	2 g.	½-1 hour	b. Ammonia (25%)	150-200 g.
(36° Bé.)	2 g.		Dissolve the chromate a,	and add b.
			Treat wood with solution,	
Cool, at 50-60°	U., add:	,	with a hard brush (horse-h	air), option.
Waterglass (30° Alcohol, Denatur	De.)	8 g.	ally a thin polish.	,, .
account, Dentur	ru.	2 g.		
T			Wood Veneer Adhe	esive
	nation Glue		U. S. Patent 1,964	
Casein	15-	-20 g.	Casein	1 oz.
Ammonia (sp. g. Water	. 0.910)8-	-16 cc.	Ammonium Sulphocyanate	2 oz.
water	77-	-64 cc.	Paraformaldehyde	.02-0.4 oz.
	_	į	Water sufficient to make i	Auid.
"Past	el'' Glue		This will remain fluid	for several
Casein		25 g.	hours at ordinary temperat	ure. Coagu-
Ammonia (0.910))	20 cc.	lates on heating to give stro	ng bond.
		•		-

Cement for Filling Cracks in Wood Consists of a mixture of 150 parts lin-

seed oil, 30 parts varnish, 40 parts wax, 30 parts gypsum, 750 parts pigment.

(Note: Generally, wax is an objectionable constituent, from the standpoints of lessening adherence within the crevices and lack of cohesion of finishing coatings applied over such filled areas. Preferable material would be the present well-known plastic wood and wood doughs which are pyroxylin-base products utilizing wood flour. Representative composition (U. S. Patent 1,838,618) is Cellu-Castor Oil 3 lbs., Ester Gum 8 lbs., Wood Flour 26 lbs.; and if pigmenta-tion may be desired, as follows—Celluloid 10, Ester 7, Castor 4, Acctone 15, Benzol 15, Alcohol 5, Wood Flour 24, China Clay 20.

Cheaper materials more popular with painters and decorators are the Water Putties in dry powder form; they are used for filling cracks and holes in wood trim, also for filling the spaces between flooring in both old and new floors. When thoroughly dry the applied putty has no tendency to shrink or crack. One product on the market for years is composed of 10 parts Quartz Silica, 2 parts Plaster of Paris, 1½ parts Dextrin. Pulverized Gum Arabic could be substituted for the dextrine and effect greater hardness; and addition of about one half part of wood flour or fine sawdust would enhance the toughness of the putty. For using, only enough water is mixed with the putty powder to the consistency of regular commercial putty).

Wood Vanour Glue

wood veneer dide	
Blood Albumen	40 g.
Casein	12 g.
Slaked Lime	6 g.
Sodium Fluosilicate	2 g.
Wood Meal	40 g.

Apply the adhesive by putting it on both sides of the middle piece of wood. If the adhesive is just too viscous, homogenize the adhesive layer. The wood pieces are put together, then pass through drying chambers at 90-95° C., under a pressure of 12-18 kg. per cc. until the albumen is congulated.

Sealing Preparation for Wine-Barrels Vaseline (40-42° C.) or so-called "Traction-Paraffine" (42-44° C.) 98-6 98-98.5 g Tallow, Hard Fat or Palm

Oil

Impregnating "Green" Wood Austrian Patent 142,431

Cover with the following paste and allow to remain until dry.

Sodium Fluoride 80 lb. Sodium Dinitrophenolate 15 lb. Kieselguhr 5 lb. Water sufficient to make paste.

Gum Arabic Glue

Gum Arabie	15-20	g.
Lame Water, Saturated	10 - 20	cc.
Glycerin	1- 3	g.
Water	74 - 27	cc.

Mucilage

Gum Arabic, Amber Sorts Water	100 150	
Heat and stir until dissolved.		

Strain and add	
Oil of Cloves	5 oz.
Oil of Wintergreen	5 oz.
Salicylic Acid	5 oz.

Photo Paste

Gum Arabic	30 g.
Saturated Lime Water	15 ee.
2% Tragacanth Solution	10 ec.
Water	45 cc.

Cold Water Paste Australian Patent 8259

Wheat Flour		8	oz.
Alum		1	02.
Water		8	oz.
Mix till smooth;	evaporate	till	dry;

Pasting Paper on Metal Surface

- 1. Clean off grease with hot soda solution.
- 2. Roughen with emery paper.
- 3. Prepare glue:

١.	Water	4	kg.
	Calcium Chloride		kg.
١.	Bone Glue	1-2	

Dissolve a, then swell b in the solution for 24-30 hours; heat on water bath to obtain solution.

Moldex or other preservative 0.1-0.2%.

Vegetable Mucilage

a. Water (Above 16° C.) 200 1. Potato-Starch b. Caustic Soda (35° Bé.) 28 kg.

Stir a to dispersion, sift, add slowly b under stirring, until glassy. Keep temperature low if thick mucilage is wished.

Higher temperature yields more fluid glues.

Library Adhesive Paste

a.	(42–44° Bé.)	70	kg.
b .	Water, Boiling Borax	20 8	l. kg.
o.	Caustic Soda (40° B6.)	2-3	kg.

d. Sulphurous Acid (5° B6.) 0.5 kg.
c. Formalin 0.5 kg.
Add b. c. d. c. in the given order se

Add b, c, d, e, in the given order separately to a, stirring strongly. When ready, dye with a little burnt sugar color.

Carton Glue

Dextrin, Light	100	g.) dissolve
Borax Solution (10%)	70	$\left. egin{align*} g. \end{array} ight. \left. egin{align*} dissolve \\ hot \end{array} ight.$
Caustic Soda		add when
(40° Bé.)	5	g. S cool
Let stand several	days.	

Cardboard Glues

1. Casein	13 g.
Trisodium Phosphate	1 g.
Ammonia (0.91)	2 g.
Water	85 g.
2. Casein	10 g.
Borax	2 g.
Glucose	2 g.
Waterglass (30° Bé.)	15 g.
Water	71 g.
	Ü

Padding Glue

1. Glue (Nat. Assoc.		
8-10 Grade)	10 lb.	
2. Glycerin	10 lb.	
3. Water	12 lb. 2 oz.	
4. Zinc Oxide	1 lb. 3 oz.	
5. Beta Naphthol	1/4 oz.	
6. Methyl Salicylate	1 oz.	
200 0 2 4 12		

Mix 2 and 4, then add 5 and 3, and then 1. Let stand over night, warm and, str until uniform; add 6 and pack.

In hot humid weather this glue may set too slowly. This may be corrected by

- a. Using a higher grade of glue, or b. Using less glycerin (which will, of
- course, reduce flexibility), or c. Dusting surface after partial dry-
 - . Dusting surface after partial drying with talc or precipitated chalk.

Tabbing Compound U. S. Patent 1,966,389

775 parts of uncoagulated vulcanized latex, containing 40 to 42% by weight of

total solids constitutes the first ingre-

The second ingredient is prepared by dissolving 50 parts of casen in about 150 parts of distilled water (preferably with the addition of an alkali which may be caustic soda, alkaline sodium salts or ammonia).

ammonia).
Third, 50 parts of egg albumen are dissolved in about 200 parts of water to produce a highly viscous solution.

produce a lightly viscous solution.
A fourth component is made by adding 125 parts of a 2% ammonia solution, to 5 parts of dried wood fibre and 5 parts of cellulose flocks (or other fibrous material) and the mixture is stirred until a substantially uniform suspension is produced. A small amount of a decodorant composition such as oil of wintergreen can also be added at this point if desired.

The casein solution and the egg albumen solution are then added slowly with constant stirring to the vulcanized uncongulated latex, and the stirring is continued until a uniform or homogeneous mass is produced. If desired, suitable coloring materials can be added at this stage and can be thoroughly stirred in.

The ammoniacal liquor containing the fibrous material "fourth component" is then added and the entire mixture thoroughly stirred or churned in order to produce a uniform mixture. This mixture is then ready to be used for tabbing, or it can be simply canned and used at any

subsequent time.

For tabbing, the paper is jogged if desired to give a substantially smooth surface of edges, to which one coat of the material is brushed on rapidly. Then after five or ten minutes a second coat is preferably applied. This second coat can be daubed on heavily, and quickly brushed down to a smooth coating. The composition will dry firm and the exposed surface will be substantially free from tackiness in about half an hour or sometimes twenty or twenty-five minutes, depending upon atmospheric conditions. The complete strength of the composition is however not developed for several hours after application. If desired, the tablets can be allowed to stand quiet for several hours, until substantially the maximum strength has developed. The surface can be finally dusted over with a suitable pulverulent material, such as talc powder if desired, although ordinarily this will not be found necessary, since the composition after drying does not stick to other surfaces with which it comes into contact, at least to an objectionable degree.

The brushes or the like used in applying the composition can be readily cleaned by being washed in water, and any of the material which gets onto the hands of the user can be readily washed off with water.

In case the solution becomes too thick, it can be diluted with soft water (preferably rain water or distilled water). Hard water would be injurious to the

compound.

I abol Com

Label Gum			
Formula No. 1—Fluid	l		
Gum Arabic Saturated Lime Water Glycerin Water No. 2—Less Fluid Gum Arabic Aluminum Sulphate Crystals Glycerin	30 g. 15 cc. 1 g. 51 cc. 35 g. 2 g. 2 g.		
Water No. 3-Viscous	61 cc.		
Gum Arabic Aluminum Sulphate Crystals 2% Tragacanth Solution Water	30 g. 2 g. 20 ec. 48 cc.		
Label Glue			
Formula No. 1 Casein Ammonia (sp. g. 0.910) 30% Rosin Soap Water No. 2	20 g. 16 cc. 5 g. 59 cc.		
Water-Resistant Casein Ammonia (0.910)	20 oz. 5 oz.		
Waterglass (30° Bé.) Water	6 oz. 70 oz.		
Library Mucilage			
Formula No. 1-Fluid			
Gum Arabic Saturated Lime Water Glycerin Water	25 g. 15 cc. 1 g. 59 cc.		
No. 2—Less Fluid Gum Arabic Lime Water, Saturated Glycerin Water No. 3—Viscous	40 g. 20 cc. 2 g. 38 cc.		
Gum Arabic Aluminum Sulphate Crystals 2% Tragacanth Solution Water	20 g. 2 g. 15 cc. 63 cc.		

Paper	Mucllage
-------	----------

a.	Dextrin, Middle Pale	50 oz.
	Water	50 oz.
b.	Sodium Bisulphite	0.5 oz.
	Borax	1.0 oz.
	Camphor	a grain

Stir cold until lump-free, warm until the mucilage is formed. Add b for deodorizing and preservation.

Adhesive for "Gunning" Papers

Gum Arabic	30 g.
Saturated Lime Water	15 cc.
Glycerin	2 g.
26 Tragacanth Solution	5 cc.
Water	48 ec.

Paper Bag Glue

Casein Borax Venice Turpentino	22 g. 3 g. 3 g.
Water	72 ec.
701	/ 11 1

The casein has to be treated (swelled) at 50-70° C. When treating with ammonia, heat up higher at the end to evaporate the excess. Moldex or other good preservative is to be added after the alkaline treatment in proportions of about 18-25 ounces per 100 gallons. If too viscous or too thin, add or evaporate water.

Let stand to clear up.

Carton Glue

Casem Caustic Soda (36° 30% Rosin Soap Water	25 Bé.) 0.5 or 1.7 10 64.5-63.3	g.
matti	01.0-	

Waterproof Adhesive U. S. Patent 1,965,778

Formula No. 1	
Casein	100 lb.
Water	225 lb.
**Wax Solution	3 lb.
No. 2	
Vegetable Protein Glue	100 lb.
Water	205 11.

Water	325 lb.
*Wax Solution	3 lb.
* Consists of	
Carbon Bisulphide	8 lb.
Carbon Tetrachloride	8 lb.
Paraffin Wax	1 lb.

Non-Caking Dextrin Adhesive French Patent 783,963

Dry adhesives having a basis of dextrin which dissolve in cold water without caking are made by heating dextrin to

about 80° C. for 1/2 hou		Celluloid Ceme	nts
of a polyhydric alcohol, e.g., glycol.		Formula No.	1
		Pyroxylin	● 200 g.
		Camphor	40 g.
Mucilage for Paper, 1		Gum Elemi	8 g.
Matter		Amyl Acetate	2600 cc.
a. Soft Water	35 g.	Acetone	500 cc.
Sugar	1 g.	Methanol	400 cc.
Wheat Starch	3 g.	No. 2	
Warm and stir until g	dassy.	Celluloid Shavings	240 g.
b. 19 parts of a 20-	25% guin arabic	Gum Elemi	8 g.
solution.		Acetone	500 cc.
Solution b is added	to a when the	Methanol	1500 сс.
starch has become "gl		Amyl Acetate	1500 cc.
with phenol or oil of cl	oves.	No. 3	***
		Pyroxylin	160 g.
Commend Intellector	Desage Office	Camphor Methanol	40 g. 2100 cc.
Gummed Labels fo	r Brass, Tin	Fusel Oil	1400 cc.
Moisten with:	0.4	Castor Oil	280 cc.
Acetic Acid	8 fl. oz.	No. 4	200 00.
Glycerin Water	2 fl. oz. 6 fl. oz.	Celluloid Shavings	40 g.
water	0 11. 02.	Gum Elemi	8 g.
		Benzol	1000 cc.
U. S. Postage St	tamp Glue	Amyl Acetato	1000 cc.
Gum Arabic	1 lb.	Methanol	800 cc.
Starch	1 lb.	Acctone	600 cc.
Sugar	4 lb.	No. 5	
Distilled Water suffic	cient to give de-	Pyroxylin	150 g.
sired consistency.	cicite to give de-	Camphor	40 g.
21104 0040120111031		Methanol	2525 cc.
		Amyl Acetate	1260 cc.
Adhesive for Wa	xed Papers	0 11 0 11 11	
Formula N	To. 1	Cement for Safety "Me	
Thickened Spiri	t-Laconer	The formula below was	
or		pecially for safety films ar	id acetate type
Acetyl Cellulose	Solution	of transparent sheeting.	
No. 2		Cellulose Acetate	4 oz.
Rosin	60 g.	Tri-Phenyl Phosphate	2 oz. 60 oz.
Mustic	10 g.	Acetone Di-Acetone Alcohol	9 oz.
Sandarac	20 g.	Benzol	15 oz.
Ether	5 g.	Methanol	10 oz.
Alcohol	75–100 g.	The cellulose acetate of	
No. 3		film quality is preferre	
Chromium G	lelatin	washed safety movie film	free from the
or		gelatin coating, or other	source of re-
Canada Ba	lsam	claimed cellulose acetate	may be used.
No. 4		Instead of tri-phenyl pho	sphate plasti-
a. Cologne Glue (or		cizers of the toluene sulp	
Gelatin)	100 g.	such as the Santicizers may	be used.
b. Acetic Acid, Dilute	e 200 g.		•
c. Potassium Bichron		Movie Film Cen	ent
Soak a in b, then d	-	This composition is effect	tive on either
bath, add c.		the inflammable or safety	type films. In
No. 5		using this cement it is scrape off the gelatin co	preferable to
Alcohol	100 g.	scrape off the gelatin co	ating with a
Ether	5 g.	knife or steel wool.	
Rosin	60-70 g.	Cellulose Nitrate	4 oz.
Sandarac	20 g.	Tri-Cresyl Phosphate	2 oz.
Mastic	10 g.	Ethyl Acetate	55 oz.

Butyl Acetate	14 oz.	Mailing Tube Adhese	va .
Benzol	15 oz.	Glue, Ground Animal	40 oz.
Methanol	10 oz.	Water	54.7 oz.
The cellulose nitrate may	consist of a	Nitrie Acid	5.0 oz.
good grade of high viscosity	nitro-cotton	Phenol	0.3 oz.
or clean new celluloid scrap	or nitrate	Combras 6 ((T)	Lalia alaa 11
movie film with the gelatin	coatings re-	Scaling of "Transparit," "I or "Cellophane" Pack	ages
moved. If new cellulose ni used, the tri-cresyl phosphate		a. Methyl Acetato	80 cc.
duced about one half. The		Ethyl Lactate	20 cc.
mixed together in the above		b. Collodion-Wool or washed	film scrap,
by weight and the cellulose ni		as much as necessary to	give a vis-
		cous solution (like 30-31)	glycerin)
Pyroxylin Cement	;		
Celluloid Scrap	40 g.	"Cellophane Adhesive	,''
Amyl Acetate	350 cc.	Arabie, Gum	16.5 oz.
Wood Alcohol	100 cc.	Glycerin	20.5 oz.
Ethyl Alcohol, Denatured	50 cc.	Glyceryl Bori-borate	9.0 oz.
Gum Elemi	15 g.	Formaldehyde	4.5 oz.
Methyl Cellulose Adh	carre	Cardboard and Nitrocellulo	so Sheet
Methyl Cellulose	1 lb.	Cement	
Water	40-60 lb.	U. S. Patent, 1,969,4	17
Warm together and stir ur	itil uniform.	Nitrocellulose	4.5 oz.
		Camphor	1.0 oz.
"Cellophane" Adhe	sivo	Acetone	30.0 oz.
U. S. Patent 1,972,		Ethyl Lactate	10.0 oz. 55.0 oz.
	-	Xylol Water	5.0 oz.
Chlorinated Polyphenyl Resin (125° C. softening			-
point)	62.5 lb.	Liquid Scaling Wax	
Dibutyl Phthalate	5.4 lb.	French Patent 751,68	33
Silica, Finely Ground	32.1 lb.	Turpentine	100 cc.
		Shellae	150 g.
Cigarette Paper Adh	esive	Zinc Oxide	30 g.
Formula No. 1		Methanol	25 сс.
Pectin Pormula No. 1	54 oz.	Mix until free from lumps.	This dries
Bone Glue, Liquid	13.5 oz.	in air after applying.	
Bone Glue, Solid	13.5 ez.		
Dextrin	19 oz.	Elastic Scaling Wax	(
No. 2		Rubber Latex (60%)	165 oz.
Pectin	60.5 oz.	Shellac	12 oz.
Bone Glue, Fluid	16.5 oz.	Warm together with stirring	g until all
Bone Glue, Solid	6.6 oz.	moisture is driven off.	•
Dextrin	12.5 oz. 4.0 oz.		
Rye Flour No. 3	4.0 02.	De Khotinsky Type Laborate	rv Cement
Pectin	50 oz.	Improved Type	•
Bone Glue, Solid	10 oz.	1	100
Dextrin	10 oz.	Shellac, Flake	100 g.
Rye Flour	5 oz.	*Plasticizing Solvent 15	
In the above formulae a	dd sufficient	Heat the solvent to 120° C., stir in the shellac flakes.	When the
water to make a mucilage of		shellac is thoroughly dissolve	
sistency.		mixture homogeneous, cool al	
		the mixture pours with difficu	lty. Imme-
Primer for Wall Paper		diately pour into long tin mol	ds of about
U. S. Patent 2,005,		one-half inch square cross se	ction which
Sodium Silicate	50 oz.	have previously been treated l	lightly with
Water	44 oz.	petrolatum.	
Copper Sulphate (121/2%	80- 6 oz.	* As a "plasticizing solvent" been widely recommended, but	pine tar has
lution)	0 024	i been mineral recommender, put	

since the excessive amount of 60 to 100 grams is required. The oil distilled from white-pine tar over the range of 200° to 325° O. is much butter, yielding a tougher cement. Wood creosote or similar inixtures of substances like guaiacoi, cresol and other low-melting, high-boiling phenois may be used, also trimethylene glycol or other slightly oxygenated organic solvents of high boiling point The range of 15 to 30 grams approximately covers the variations of hardness commonly desired.

"Boltwood Wax"

(For cementing physical instruments)

(Tor cementing Improm	, and the distriction of
Shellac	40 g.
Rosin	72 g.
Venice Turpentine	8 g.
Beeswax	60 g.
Tale, Dry	16 g.
Tin Oxide, Dry	16 g.

Melt the rosin, add the shellac. Heat to 200° C, add the Venice turpentine and becswax. Heat the mixture strongly with stirring until it ignites spontaneously. Let it burn until the total mass has shrunk to about 40% of its original weight, then add the tale and tin oxide. This gives a tough, smooth, waxy cement more easily handled on certain delicate instruments than the de Khotinsky type cement.

Leather Sole Cement

Nitrocellulose	22.5 g.
Alcohol	22.5 g.
Benzol	31.1 g.
Ethyl Acetate	9.5 g.
Camphor	1.1 g.
Acetone Oil	0.09 g.
Castor Oil	0.09 g.

Cement for Leather or Leather on

Rubber		
Gutta-Percha	21.6	oz.
Carbon Bisulphide	17.7	υZ.
Benzene	2.9	oz.
Turpentine Oil	23.5	07.
Asphalt	34.3	OZ.

Leather Cement

Celluloid	11.9	oz.
Naphthalene	1.2	
Acetone	67.1	oz.

Cement for Stone and Leather, Porcelain and Leather, Glass and Leather

Crude Rubber		9.1	oz.
Heavy Benzine		45.5	
Japan Wax		13.6	
Colophony		31.8	0 Z.

Concentrated Rubber Cement German Patent 599,405

a.	Caoutehouc Benzol	•	10 g. 90 g.
ħ.	Nitrie Acid	(52.77%)	1 g.

a gives after 24 hours stirring a homogeneous puste, which is depolymerized by adding b. When paste is dissolved, stop reaction by adding barium enrhonate. Treat then with antimony trichloride or phthalic acid.

Rubber Cement

(Will firmly fasten rubber to almost any substance)

substance)		
India Rubber (finely		
chopped)	100	oz.
Rosin	15	oz.
Shellac	10	oz.
Carbon Disulphide, sufficient	to di	ssolve

Softening Hardened Shoe Adhesive German Patent 605,725

Cellulose nitrate adhesives used in shoe cements are softened by the following:
Pyroxylin (1100 second) 62 oz.
Alcohol 26 oz.

Alcohol 26 oz. Acetone 450 oz. α-Propylene Oxide 225 oz. Shoe Repair Cement

U. S. Patent 2,004,059 Crepe Rubber 6 lb. Rosin 2½ lb. Accelerator 114 lb.

Accelerator Benzene	11/2	
Porous Leather	Sealer	

Shellac	14	lb.
Rosin	1	lb.
Alcohol	5	gal.
Butyl Alcohol		gal.
Castor Oil	4	OZ.

Leather Belt Cement

a. Glue, Hido	50 g.
b. Water	200 g.
Soak a in b, pour excess	water off, and
melt the soaked a with:	

c. Glycerin 2% Potassium Bichromate 2%

When cooled, pour into oiled metallic forms; pack the gelatinous product at once into grease-proof paper.

once into grease-proof paper.

Apply on roughed surface, while the sharpened ends are pressed together for 6 to 10 hours.

Belting	Cement

Hide Glue	2¼ lb.
Water	2¼ lb.
Glycerin	9 oz.
Carbolic Acid	3 ₁₆ oz.
To use, melt and appl-	y hot to the
eather belt and place the	
pressure until the glue is the	horoughly set.

Canvas Awning Cement U. S. Patent 2,011,218

Rubber Latex	10	oz.
Varnish	1	OZ.
Citronella Oıl	1/100	υz.
Nigrosine B Solution	1/100	

Textile Glue

(for Doubling of Cloth, Shirting, Drill) 15 oz. Soft Soap, Pure 5-10 oz. 2 oz. Borax Water 75 oz.

Warm and stir together.

Jute or Burlap Sheet Binder British Patent 412,498

Gilsonite	11 lb.
Asphalt, Petroleum	23 lb.
Naphtha, Petroleum	35 lb.
Mineral Silicate Filler	15 lb.
Asbestos, Fibrous	15 lb.
Linseed Oil	2 lb.

Upholsterer's Paste

Prepare a

a. Calcium Chloride Solution (25° Bé.)

cleared by pouring off solution from settled dirt, and add 160 kg.

b. Potato-Starch Water

100 kg. 100 l.

(Heated to 60-65° C.) This glue has a good binding power, but dries very slowly and is hygroscopic.

Fine Bookbinder's Paste T):-- 1 . . .

Dissoive in	
Water, Boiling	100 l.
Trisodium Phosphate	15 kg.
{ Borax	2.5 kg.
or { Alum	10 kg.
and add with stirring, a	solution of:
Water, Cold	120 l.
Starch	50 kg.
Warm until fluid.	

Upholsterer's and Bookbinder's Paste

a. Potato-Starch 140 1. Water, Cold

b. Caustic Potash (50° Bé.) 6 kg. 15 kg. Sodium Silicate Water, Cold 50 1.

c. Acid to neutralize to weak alkalimity d. Rosin Soap, Warm Flind 5 kg.

Stir a till smooth, warm and stir with b to form a mucilage. Stir 3/4 to 1 hour more, add c, then d, and stir slowly.

Bookbinder's Paste

a.	Rye or Wheat Flour	100 kg.
	Water, 25° C.	200 1,
b.	Caustie Soda (35° Bé.)) 24 kg.
c.	Nitric Acid	until neutral

d. Alum, Cold Saturated Solution

20 kg.

Stir a to dispersion, treat mildly with b, neutralize with c, and add d.

Adhesive Paste for Rubber-Cloth on Cardboard

a. Gutta Percha, Finely Cut 18 g. 20 g. Carbon Disulphide 10 ğ. Benzene 10 g. Turpentine Oil b. Colophony 42 g.

a is mixed and soaked several days, then add b with gentle warming.

Mending China, Pottery and Casts

Save all the pieces of the broken article and store where the edges will keep clean until the repair is made. If the edges become soiled they should be washed clean and allowed to dry. The edges may be sanded lightly if necessary to remove the soil. The worker should know where each piece belongs before the work is begun. Small pieces should he cemented together previous to the main repair. A sand box is convenient to hold pieces upright while making the repair leaving both hands free for the work. It is made by putting 8 mehes of clean sand in a convenient sized box.

Have at hand the cement, rubber bands, a bowl of warm water, tissue and soft rags. One rag should be reserved for wiping the fingers. Do not work with sticky fingers. Be accurate. If some part is not true after having been put together, soak until the cement is dissolved, wash the edges and begin over. Warm water will dissolve plaster or whiting cement and turpentine or alcohol will dissolve others.

ľ

The most durable cement is pure white lead ground in linseed oil, so thick that it will barely spread smooth with a knife. After drying thoroughly (about three months) it makes a seam which is practically indestructible but the mend is very conspicuous.

A less conspicuous cement is made of beaten egg white and sifted whiting or plaster of Paris. A small amount should be mixed at a time as it hardens quickly. In some cases it is just as satisfactory to brush the edges with beaten egg white and dust well with sifted plaster tied loosely in double mosquito netting. The pieces should be fitted together at once and held in place by rubber bands (placed lengthwise, crosswise and diagonally) wrapped loosely in tissue paper and buried in a sand box. Care should be taken that the break lies so that the weight of the sand will hold it together. Leave it in the box at least 24 hours. After a week the superfluous plaster may be scraped away.

Sometimes the rubber bands will not hold the pieces true on a stemmed article, a vase or a jug. In this case string six bands of the same size and strength upon a piece of tape. Tie the tape around the neck or base of the article before beginning the gluing. After the parts are joined slip another tape through the bands and tie above the fracture. The bands pulling in unison will hold the break together. The pressure on all mended fractures should be great enough to force out the tiny air bubbles which otherwise reflect light making the seam conspicuous.

Universal Putty for Wood, Stone, Glass, Porcelain

(Dries after 24-30 hours)

a. Alabaster Gypsum 4 oz.
Gum Arabic 1 oz.
b. Cold Borax Solution, Saturated.
Stir until pasty.

Preserve Jar Scaling Wax Washes off easily with hot water. Paraffin Wax 35 g. Trihydroxyethylamine Stearate 3 g.

Paraffin Bottle Cap Adhesive U. S. Patent 1,964,380

Chicle 1 oz.
Dammar 1 oz.
Petrolatum, Liquid ½ oz.
Warm and stir until homogeneous.

Bottle-Cap Varnish

Dissolve 2 oz. of red Sealing-wax in 5 oz. of denatured alcohol.

Seal for Bottles

Beeswax	5 g.
Carnauba Wax	1 g.
Paraffin	1 g.
Minium	5 g.
Whiting	2 g.

To Seal Glass Tubing to Iron Tubing Grand the ends you wish to join to a tapered fit and then seal by fusing with silver chlorade,

Cement for Vacuum Tubes

Marble Flour	85	oz.
Shellac	10	oz.
Rosin	5	υz.
Phenol Formaldehyde Resin	25	oz.

Glass to Metal Scals

Formula No. 1

ron	37 oz.
Vickel	30 oz.
Cobalt	25 oz.
'hromium	8 oz.

The above is suitable for use with leadglass.

No. 2

ΩU	
20	lb.
	lb.

Suitable for use with Corning glasses.

Safety Glass Adhesive U. S. Patent 2,009,029

Formula No. 1

A small portion of casein is heated in an open vessel with twice its weight of glycerol and 1.0% by weight sodium hydroxide (based on the casein). The temperature is brought gradually to 150-165° C. over a period of 15 minutes with continual stirring, and then held at this point for an additional 30 minutes. This product is a clear liquid at 100° but rubbery and very slightly opaque on cooling to room temperature. This material while hot may be pressed between two hot pieces of glass until air bubbles disappear. On cooling a piece of sandwich

glass is obtained in which the glass plates are firmly held together.

No. 2

Fourteen and nine-tenths (14.9) parts glycerol, 35.1 parts phthalic anhydride and 10.0 grams sheet gelatin (broken into small pieces) are heated with strring in an open aluminum vessel, one hour up to 200° C. and 4 hours at 200° C., or to an acid number of 65-70. Some difficulty may be experienced in the early stages in making the bulky masses of gelatin mix with the other maternals. This resin may be used as the sandwiching material for glass, or dissolved in a solvent such as acetone and used as an adhesive or impregnating agent.

Percent Quartz					
Coefficient of Expansion					
Percent Porcelain					
Coefficient of Expansion					

The quartz cement mixtures for values of quartz between 40% to 70% usually shows the same coefficient of expansion as pire cement. The modulus of clasticity of the quartz cement mixture increases with increasing quartz content. The bending strength, however, decreases almost in proportion to the percent quartz. The impact or shock bending strength, however, is practically unaffected up to 50% quartz content.

Porcelain and metal surfaces should be given a coating of a good elastic varnish before cementing. The cement should be allowed to harden in a steam chamber or, at least, be kept thoroughly wet for the first forty-eight hours.

Another good porcelain cement is the usual litherge glycerin cement. This should be made in a ratio of three parts litharge and 1 part glycerin by weight. The glycerin used should contain less than 15% water and the litharge must, as far as possible, be free of lead carbonates as they produce a porous, weak cement.

A filler of up to 40% crushed or powdered porcelain may alse be used advantageously with the litharge. All exposed surfaces of cement should be given a thoroughly protecting coating of a good grade of Glyptal or Bakelite varnish.

Litharge and glycerin ratio about 75/25 sample poured in a 25 mm. diameter glass tube hardens to a solid mass in less than 24 hours, but on further drying gives off additional moisture thereby slightly decreasing its dimensions so that it can be pushed out of tube. Swells

Mastic Scal for Oil Drums German Patent 613.748

Aluminum Powder	30 kg.
Nitrocellulose	14 kg.
Butyl Acetate	21 kg.
Ether	35 kg

Glass Electrical Cements

To offset the greater thermal coefficient of expansion of ordinary cement (11.5 × 10⁻⁶) against that of porcelain (4.5 × 10⁻⁶) a mixture of cement and powdered quartz or cement and crushed porcelain may be used. The thermal coefficient of expansion has approximately the following values:

0	20	40	70	80
11.5	10	8.5	5.5	4×10^{-6}
0	20	40	60	80
11.5	10.5	9	7.5	6×10^{6}

on moist days sufficiently to firmly hold sample in glass tube. It is now adhering to glass. Under the microscope it shows a fairly dense even mass with numerous minute air-bubbles which appear to be conted with a shiny scale. Cracks when heat is locally applied and apparent traces of glycerin start to burn with a slow glowing, causing bubbles to be formed. Mechanically very rigid and strong, water absorption in 14 hours—1.6% by weight.

2.

Equal parts litharge and crushed porcelain plus glycerin to make a good flowing cement. Hardens in less than 24 hours, forms a hard solid body which cannot be moved in glass tube but under the microscope shows somewhat more porous than No. 1, especially around the coarser grains of crushed porcelain. Mechanically rigid and strong.

2

Glens Falls Cement Company iron clad portland cement and water. Cement poured in 25 mm, diameter glass tubes, hardens in less than 24 hours but 7 days is recommended by the manufacturer to give it full strength. One test tube was kept under water for the first 48 hours according to the recommendation of the manufacturer and one tube air dried only. The air dried cement could be hammered out of glass tube and under the incroscope showed minute air bubbles imbedded in the solid material. The sample set under water showed a very dense homogeneous body composed of minute bright crystals imbedded in a

mass of various dull colored material. The sample set under water showed considerable more strength and toughness than the air dried absorption in 14 hours -8.8% by weight.

50% "iron clad" portland cement, 50% crushed porcelain. Sufficient water to readily pour sample set under water for the first 48 hours and allowed 6 days for air hardening. This sample gave a hard tough body of high mechanical strength.
Under the microscope it showed the porcelain particles very densely imbedded in the material and traces of air bubbles could only be found around the larger porcelain grains. It appears to be a very promising cement for porcelain cementing. Number 4 very closely resembles the so-called "Teleo" Cement patented by the porcelain factory Treiberg in Thyringen, Germany, and consisting of portland cement and crushed quartz glass. This cement was developed with a view of obtaining a cement of approximately the same temperature expansion as that of porcelain. This is obtained by mixing a sufficient quantity of crushed quartz glass with an expansion coefficient of 0.5 × 10-6 with the portland cement having an expansion of 11.5 × 10-6 to give an expansion coefficient of approxi-mately equal to that of porcelain of 4.5×15^{-6} . Further tests on the various cements are necessary in order to fully determine the mechanical properties.

Summary

The indications from the above preliminary tests, therefore, are that litharge and giveerin in a ratio of about 80,20 by weight or a mixture of 7 parts Glens Falls iron clad cement and 3 parts powdered porcelain or perhaps still better powdered quartz and water is the most suitable cement to use for bushing work.

The metal and porcelain surfaces to be given one coat of clear "Valspar" varnish to take care of the variation in expansion and all free surfaces of the cement to be given two or three coats of varnish as a protection against moisture.

To Plug Holes in Metal

Mix powdered sulphur and powdered aluminum 1-1 and pour on the metal which should be hot and clean. Then heat to melt the sulphur.

Metal Glue (for Tins, Etc.) Resin (Shellac, Sandarae) 50-100 g. Manila-Kopal, Soft 50- 0 g.

FORMULARY	
Galipot or Turpentine Thick Alcohol, Denatured Castor Oil	3 g. 100-200 g. 1 g.
Pipe Joint Lu German Patent 59	te 17,044
Tallow Mineral Oil	1 lb.
Melt together and mix	1 lb. with:
Ochre Dry Clay or Sand	1 lb. 7 lb.

Premolded Expansion Joint Chinawood Oil, Polymerized Bitumen 85 lb. Mineral Filler 10 lb.

Sulphur Thiokol Cements Formula No. 1

Sulphur Thiokol Sand		58.8 lb. 1.2 lb. 40.0 lb.
Sulphur Thiokol Sand Carbon Black	No. 2	58.8 lb. 1.2 lb. 38.0 lb. 2.0 lb.

Refractory Cement U. S. Patent 1,952,119

	440
Magnesium Oxide, Powdere	ed
(Deadburned) (Fused)	50 lb.
Zircon Sand	15 lb.
60-mesh	25 lb.
300-mesh Sodium Silicate (d. 1.3)	30 33
make paste.	samelent to

High Temperature Luting Compound Alumina 50 lb. Magnesia. 25 lb. Kaolin 25 lb. Sodium Silicate sufficient to bring to a working consistency.

Nitric Acid Resistant Putty

THE TENT TENTAL	11 Putty
White Asbestos Powder	20 parts
Blue Asbestos Fiber	10 parts
China Clay	10 parts
Linseed Oil	20 parts

A cement for nitric acid plants contains:

Blue Asbestos Powder, and Sodium Silicate 1.5 Tw.

Asbestos Binder U. S. Patent 2,010,224

Shellac 48 oz. Dicyandiamide 2 oz.

Heat together and stir until uniform.

Acid-Proof Dental Cement

Make a concentrated solution of silicate of soda and form a paste with powdered glass. Invaluable where a luting is required to resist the action of acid fumes.

Dental Cement British Patent 430,624

Lithium Phosphate	1/2 oz.
Phosphoric Acid	5 07.
Zinc Phosphate	1/2 OZ.
Aluminum Phosphato	1/3 oz.
The above is added to a	ground porce-

lain of following composition:
Alumina 30-50 oz.

Feldspar 10-20 oz. Sand 25-40 oz. Zinc Oxide 1-10 oz.

Boiler Lagging

A splendid boiler lagging can be made by the following formula and applied direct to the boiler with a trowel, or molded into sections or blocks of suitable size and then dried and applied in the form of the usual sectional lagging:

- 200 lb. spent Carbide Residue, drained to a soft putty consist-
- 100 lb. Asbestos Fiber or Asbestos
 Fiber and Magnesia. (Old lagging properly ground will be satisfactory.)
- 3. 50 lb. Fine Dry Pine Sawdust.

Mix 2 and 3, then add 1 and mix thoroughly. If too dry add a small quantity of water. If onk or wet sawdust is used, quantity should be increased in the same proportion as the difference in weight per cubic foot.

It has also been found that carbido readue mixed with equal parts of Fuller's Earth will produce a good heat insulator for small furnaces.

Silicate Cements

Composition

Methods

Remarks

Silicate of Soda

Apply to porous surface and wash with dilute sulphusic acid after setting

Silicate of Sods and Asbestos Fiber

Silicate of Soda and Silica or Clay

Silicate of Sods and Whiting

Silicate of Soda and Diatomaceous Earth

Silicate of Soda and Portland Cement

Silicate of Soda and Zinc Oxide, with or without added Clay

Silicate of Soda and Sawdust or Wood Flour

Silicate of Soda and Copper Powder Mix to paste and wash with dilute sulphuric acid to develop acidresistance after setting.

Mix to paste and wash with dilute sulphuric acid to develop acid-resistance after setting

Mix to paste and wash with dilute sulphuric acid to develop acid resistance after setting

Mix to paste and wash with dilute sulphuric acid to develop acid-resistance after setting

Very quick setting; make only as needed

Portland cement may be added

"Acid-proofing of wood,

unglazed tile, etc.

General acid-proof coment and lute; also used for setting acidproof bricks, etc. Acid-proof and refractory

For setting acid-proof tiles; waterproof

Used as a binder in abrasive wheels; waterresistant

Strong bond; water-resistant; also resistant to weak acid

For protecting spots during case hardening

Silicate Cements—Continued			
Composition	Methods	Remarks	
Silicate of Soda and Ba- rytes Flour	Make to a stiff paste	Resists wet chlorine	
Silicate of Soda and Duriron Dust	Make to a stiff paste	Used for temporary re- pairs of Duriron	
Silicate of Soda and Sil- ica Flour and Sodium Fluosilicate	Make to a stiff paste	Used for temporary re- pairs of Duriron	
Silicate of Soda and 20 Manganese Dioxide; 20 Zinc Oxide; 10 Kiesel- guhr; 3 Graphite	Make to a stiff paste	Used for repair of metal parts; becomes highly acid resistant on setting.	
	Glycerol-Litharge Cements		
Composition	Methods	Remarks	
a. Glycerol and Litharge	Mix to a paste and apply promptly; varying the proportions, changes characteristics	Proportions vary; addi- tion of water to gly- cerol hastens setting (2 water to 5 glycerol	
b. (a) plus Whiting	Slower setting than straight cement	sets in 10 minutes)	
c. (a) plus Silica	Slower setting than straight cement		
d. (a) plus Ferric Oxide	Slower setting than straight cement		
e. 1 part Litharge; 1 part Silica; 1 part Portland Cement, Gly- cerol and Silicate of Soda (diluted)	Addition of silicate con- trols setting time	Sulphite digester linings; dilute sulphuric acid	
f. 1 part Litharge; 1.5 parts Silica; 1.5 parts Portland Cement; Gly- cerol and more Silicate	Mix to a putty consist- ency	For sulphur dioxide gas (wet); resists hot so- lutions	
g. 5 parts Litharge; 3 parts Silica; 2 parts Quartz Flour; c.p. Glycerol	Mix to a putty consist- ency	For sulphur dioxide gas (wet); resists hot solu- tions	
h. Glycerol and Litharge plus Graphite	Mix to a putty consist- ency	Used on pipe joints which can be taken apart easily	
i. Glycerol and Red Lead	Mix to a putty consist- ency	Acid-resistant joints in iron; sets hard	
	Miscellaneous Cements		
Composition	Methods	Remarks	
Iron Filings (100); Ammonium Chloride (1); water	Mix to a thick paste	Used to repair cast iron, etc.; resistant to heat but not acids	
Asbestos Wicking and Rubber Cement (rub- ber dissolved in ben- sene)	Soak wicking in cement and force into joint (not too strongly)	Used as caulking on fused silica and stone- ware bell and spigot joints; proof against moisture and dilute acids; flexible	

Miscellaneous Cements-Continued

Composition	Methods	Remarks
Lead Wool	Caulk into joints	Used in the same way as poured lead joints in bell and spigot pipe
Asbestos Wicking	Used as a caulking with or without asphalt or other cement to protect it	Resists common acids except hydrofluoric
White Lead and Varnish Putty	.25 to 1.5 gal. of hard drying varnish to 100 lb. paste white lead in linseed oil	For jointing marble, stone, glass, etc.; an adhesive, slow-harden- ing cement
White Lead Paste with Read Lead	Red lead added to give the heaviest workable paste	For threaded pipe joints; can be opened
Lead Filings	Lead is filed on to pipe threads moistened with lubricating oil	Makes tight threaded joints
Red Lead in 3 parts, raw Linseed and 1 part medium Lubricating Oil	Mix to stiff paste	Adheres tenaciously to metal; remains soft and elastic; fillers may be added
Cellulose Acetate solu- tions (with or without fillers)	Applied as a scaling com- pound	General service adhesive
Cellulose Nitrate solu- tions (with or without fillers)	Applied as a scaling com- pound	General service adhesive
Rubber, Linseed Oil, As- bestos Fiber	Rubber is dissolved in hot oil and asbestos added to make a thick putty	For joints in stoneware, etc.; forms an elastic mass
Sulphur in various mix- tures	Sulphur is melted and mixed with clay, sihea, etc., to form a putty	Applied hot as a grout- ing; resists acids and alkalis
Self-vulcanizing Rubber Cement	Painted or trowelled in place	Resists both corrosion and abrasion
Numerous resin base pro- prietaries		Resist dilute acids
Synthetic resin varnishes		Resist acids and weak al- kalies
Soaps (particularly of heavy metals)	Made to a putty with lin- seed or other drying oil	Resists hydrocarbon solvents
3 lb. dry White Lead; 2 lb. White Lead in Oil; 1 lb. 85% Magnesia with Linseed Oil to make stiff putty	Laid between flanges of joints, using a lead wire as a shim	Resists hot alcohol vapors
80 lb. Litharge; 8 lb. Red Lead; 10 lb. Floc Asbestos; 1.5 gal. Lin- seed Oil	Hardens in about 7 days	Resists dilute nitric acid cold but not hot
Tar or Soft Pitch and Linseed Oil (50-50)	Applied hot	Does not harden; resists acids
Sulphur melted with Rosin Tar or Pitch	Melted in place	Resists hydrochloric acid

Miscellaneous Cements-Continued

Composition

Methods

Remarks

Shellac	(30);	Rosin
	Alcohol	(33);
Gypsun Oxide (F'erric

Finely powdered solids are mixed with an alcoholic solution of the resins

Resists petroleum oils

2 parts Scotch Glue; 7 parts Water; 1 part Glycerol For oil or gas leaks; more glycerol softens it

Non-Efflorescing Concrete

The addition of 5% Barium Carbonate to the cement prevents efflorescence.

Keying Plaster to Concrete

First secure a fast setting plaster which corresponds to Plaster of Paris, which corresponds to Pinster of Paris, moulding plaster or something similar. This plaster is mixed thin enough so it can be whipped onto the wall with a brush. After this dash coat of plaster has thoroughly set, the wall, which now have a new harders. has a rough surface, may be plastered over in the usual way with ordinary gypsum plaster.

Plaster Coment, Patching U. S. Patent 2,016,986

	4 lb.
Dry by heating below 600° C. Slaked Lime	5 lb.

Refrigerator Display Case Caulking Compound

U. S. Patent 1,974,745

Nitrocellulose	1- 7	oz.
Dibutyl Phthalate	15-60	oz.
Asbestine (Mineral)	30-90	oz.
Camphor	1/4	ψz.

Cement "Wash" Hardener Portland Cement 20 lb. Iron Filings 126 lb. Water 9 lb.

Apply with brush, mixing often.

Concrete Wash, or Finish Paint (Hard and Durable)

Slaked I	imo		1	lb.
Cement			1	lb.
		containing		salt

Colored Caulking Cement U. S. Patent 2,011,607

A cement of substantially permanent elasticity and which is adapted for application by a trowel or a grease gun consists of paracoumarone resin m.p. about 50-60° C. about 60, asbestos fiber about 20, a metallic oxide such as oxide of zinc or iron about 5 and xylol about 15%.

Pliable Glazing-Caulking Cement British Patent 398,057

Formula No. 1		
Mineral Filler	1-50	oz.
Oil	30	oz.
Asbestos Fiber	20-1	oz.
Aluminum Powder	1-30	oz.
Varmsh sufficient to make	paste.	

No. 2

Calcium Carbonate, Powdered	10.00
	12.60 oz.
Magnesium Silicate,	
Powdered	17.10 oz.
Asbestos Fiber	5.45 oz.
Soya Bean Oil	30.63 oz.
Varnish	16.22 oz.
Aluminum Powder	9.00 oz.
Naphtha, Petroleum	9.00 oz.

33 ...

Glazing Putty	
Whiting, Domestie,	
200 mesh	205 lb.
Whiting, Belgian	70 lb.
Linseed Oil, Raw	26 lb.
Japan Drier	1 lb.
Mineral Spirits	3 lb.
_	

Cement for Pestle Handles

Heat the head of the pestle until it is too hot to hold in the hand. Pour melted shellac into the hole, then take the wooden handle, wind some twine around the screw portion, and press it "home." Keep under pressure until the head of the pestle is cold.

Mortar Cement

Fuse together, in an iron vessel, equal parts of guttapercha and shellac. This forms a powerful cement. Strongly heat the edges of the broken mortar, apply a thin layer of the cement to both fractured surfaces, and put together under pressure.

Joining Stainless Steel in Knife Handles Method 1

A waterproof cement is used, made by mixing finely powdered litharge and glycerin. The glycerin should be added in an amount equal in volume to half the volume of the powdered litharge and mixed thoroughly. The end of hollow handle is filled with cement and then insert the blade. Setting time about 45 minutes. Mix only enough cement as needed as it sets quickly becoming hard and msoluble.

Method 2

The stainless steel blade is first thoroughly tinned and then soldered in place It is necessary to have all parts clean and free from scale. Solders used are either 50% tin and 50% lead or 66% tin and 34% lead. Flux used is made up of zinc chloride, commercial grade, 37 g.; glacial acetic acid 99.9%, 23 g.; hydrochloric acid (commercial), 34.5%, 40 g.

Metal Adhesive

Nitrocellulose Scrap	10 g.
Alcohol	26 g.
Ethyl Acctate	25 g.
Butyl Acctate	31 g.
Benzol	30 g.
Camphor	2 g.
To the viscous solution a	idd:
Metal Powder enough	to ''lude''
Viscosity should be log	h enough to

Rubber to Metal Cement British Patent 432,493

Paris White	40	OZ.
Rosin	3	OZ.
Dammar or Copal Gum	15	07.
Benzol	15	oz.
Naphtha	23	oz.
Rubber	1 1/2	oz.
Pyroxylin to Metal	Adhesive	

r yroxyiiu		metar	Tr.III. De . O	
Pyroxylin			6	oz.
Gelatin			7	oz.
Acetic Acid,	Gla	cial	87	oz.

Aluminum Foil to Leather or Paper Adhesive

U. S. Patent 1,925,903

Linseed Oil Fatty Acids	11.82 g.
Tung Oil	16.35 g.
Rosin	22.53 g.
Heat rapidly in aluminu	m vessel to

			to	260-265°	C.	and	add
with	stirr	ing:					

	Anhydride	32.68 g.
Glycerm		16.35 g.
Ethylene	Glycol	4.22 g.

Keep at 200-220° C. until clear; heat at 250° C. until a sample solidifies in 40 seconds at 200° C.

Take of the above resin	11 g.
and dissolve in: Acetone	11 g.
Dibutyl Phthalate Nitrocellulose "Solution"	5 g.
(1/2 second)	sufficient

Thermoplastic Cement	
Nitrocellulose Wet 5-6 sec.	8 g.
DuPont Resin RH 35	
6# cut	10 g.
Dibutyl Phthalate	4 g.
Methyl Ethyl Ketone	10 g.
Butyl Acetuto	10 g.
Toluol	58 g.
	_

Fusible Adhesive Cement U. S. Patent 1,945,803

Chlormated Naphthalene	
(Solid)	50 oz.
Ester Gum	50 oz.
Rubber Latex	5 oz.

Shellac Scaling Composition

omenas;	1,11,	
Beechwood Creosote	5	oz.
Ammonia (28%)	1	oz.
Terpincol	2	oz.

Adhesive Sealing Compound (Universum)

Mix hot beeswax and Venice turpentine 1 to 1. Proportions may be varied ac-cording to needs. Can be colored if de-sired. This is very good to temporarily attach glass to iron or wood.

"Syndetikon" (Universal Adhesive) a. Prepare Caustic Linie,

100 g. Freshly Burned 50 g. Water

Let stand to cool: pour off layer of water. Use now:

(Lime Hydrate (above) Sugar Solution (25%) 240 g.

Heat to 75° C, let stand stirring through from time to time, pour off the clear solution, of which

Lime Sugar Solution Bone Glue 60 g. are mixed to swell over night. Dissolve finally by warming up.

<u> </u>	HE CHEMIC
Acid Resisting (Coment
Fine Sand	
Short Fiber Asbestos	2 lb.
Magnesia	2 lb. 1 lb.
Sodium Silicate suffici	1 lb.
paste.	ent to make
Aquarium Cen	- ient
Litharge	3 lb.
Fine White Sand	3 lb.
Plaster of Paris	3 lb.
Mix thoroughly Then	add lingood oil
Mix thoroughly. Then sufficient to make paste,	and a small
amount of drier.	, and a sman
	_
Adhesive Fo	1
U. S. Patent 1,9	55 , 07 5
Acidify defibrinated bl with 0.5% lactic acid; n	mx with 2.3%
ammonium sulphate solution. C. for 1-3 hours; render solution mix with 8-12% gly	m; keep at 40°
C. for 1-3 hours; render s	lightly alkaline
and mix with 8-12% gly	cerin and 5%
alum or synthetic tannins.	- 70
-	-
Adhesive for Casein	Plastics
British Patent 41	11.058
Casein	
Water	1 part
Urea	1 part
	1/2-1 part
Onigh Hamlanian	D. Att.
Quick Hardening German Patent 63	11111108 13 7.19
Formula No.	
Aluminum Powder	30 g.
Nitrocellulose	14 g.
Butyl Acetate Ethyl Ether	21 cc.
Etnyl Ether	35 ec.
No. 2	
Aluminum Powder	30 g.
Ethyl Cellulose	14 g.
Benzol Valuation	33.6 cc.
Ethyl Ether	22.4 cc.
D-1 I - 1 D -	.
Red Lead Put	•
Red Lead, Dry	31 lb.
White Lead, Dry	48 lb.
Silica Prov. I	16 lb.
Raw Linseed Oil	1 gal.
	•
Slate Color Put	tty
Whiting	24 lb.
White Lead, in Oil Lampblack, Dry	70 lb.
Pour Lineau Lory	2 oz.
Raw Linseed Oil	6 lb.
	•
White Putty	
Whiting	77 lb.
White Lend in Oil	0.11

White Lead, in Oil

Raw Linseed Oil

9 lb.

Black Plastic Put	ty
"D" Asphaltum (Soft)	400 lb.
Gilsonite	100 lb.
Black Fish Oil	7 gal.
Crude Black Oil	7 gal.
Stove Distillate	70 gal.
Directions:	6

Melt the two blacks to 550° F. and hold until in complete solution, then add both oils and heat to 575° F. Cool to 450° and reduce.

The black fish oil is a very dark crude and chenp oil, unfiltered and full of stearines.

For overglazing where the lights of glass overlap, a semi-liquid coating is made by mixing into the base vehicles while lot ¾ lb. of long-fiber asbestos to each gallon.

For the plastic putty for cementing the glass to the frame, the following mixture is made in a regular pony chaser:

Base Vehicle (above) 5 gal. Stove Distillate 1½ gal. ''Asbestine'' 50 lb. Long-fiber Asbestos 5 lb.

This product is stiff and must be applied by knifing or with a small trowel.

In the cast and south cement slabs called cementiles are quite commonly used in constructing factory roofs. The joints of these tiles are first partly filled in with a non-shrinkable cement, and above this flush with the tile surface is run a waterproof expansive plastic for protection. An eastern manufacturer of cementile roofing slabs also makes the joint cement or putty. They buy large quantities of paint skins from paint manufacturers, and use this as the base material, cooking the same with an addition of fish oil, subsequently churning it with such filling material as asbestine or whiting, short asbestos fiber, and red oxide for color. The final protective is a wellknown commodity, trade name similar to "mud mud." Its salient features are: a soft but firm plasticity; a condition of slime for easy slip in trowelling; slow setting during manipulation, but later becomes surface set out of dust and dirt; retains its softness and cohesiveness within the joint, indefinitely. These features have been very well reproduced in the following formulation:

5% Leaded Zinc Oxide	24	lb.
Borate of Manganese	1/4	lb.
Spanish Red Oxide	8 -	lb.
Treated China Wood Oil	4	gal.
Sulphurized Fish Oil	4	gal.
Medium Body Gloss Oil	4	gal
"C" Asbestos Fiber	32	lb.

The prepared oil is 40 lb. of limed rosin and 20 gal. of wood oil heated to 455°F. and held there about 2 hours until very heavy—but no stringing; then reduced immediately with 50 gal. kerosene.

The above plastic is run into the tile joints with a hand-pressure caulking and glazing gun, fitted with either the stundard or the extra large caulking nozzle.

Although akin to putty but more properly termed otherwise, is that compound familiarly known by almost everyone as Litharge Glycerin Cement, which is val uable for a number of purposes for which ordinary cement and putty would be neither practicable nor desirable. Probably all readers may feel that they know how to mix this cement for usage, but those who merely combine these two in gredients really would not be doing it efficiently for best results. The cement is correctly produced by adding to a mixture of 5 parts of 95% pure glycerin and 3 parts of water, sufficient finely ground litharge to form a plastic of any required consistency. Variation in the amount of water will influence the time of setting and to an extent the general characteristics, but all modification within the range of say 1 to 3 parts of water with 5 or 6 parts of glycerm will attain satisfactory hardness. Its normal hardening time is about ten minutes, but it may be made to remain soft for a langer period by an addition of ten per cent of mert material such as silien, iron oxide, or fuller's earth. Such admixtures do not detract from the ultimate hardening or strength, but also are beneficial in preventing possible cracking. Litharge-Glycerin Cement will with stand a high degree of combined heat and moisture. A very common usage is for forming water-tight connections between iron pipes and porcelain fittings; and for cementing glass aquariums, etc. Its most conspicuous feature is its resistance to practically all acids not of full strength. It is used to good advantage in temporarily scaling leaks at seams, around the bottoms, and around flanges, etc., of storage tanks filled with varnish; these temporary repairs have held until the contents of the tanks were used when a permanent repair could be made

Marine Putty, to harden under water, may be made from the formulation here given:

Hydraulic Cement	30	lb.
Plaster of Paris	71/2	lb.
Litharge	10	lb.
Belgium Whiting	20	lb.

Lead Carbonate (Dry) 10 lb. Boiled Linseed Oil 3 gal.

On the scaboard, hydraulic cement is better known as sea-water cement. This type differs from regular Porthand cement for land construction in being darker color and construction in being darker color and containing a minimum of tra-calcium aluminate... the constituent in cement which is rapidly attacked by (salue) sea water. Whereas regular cement contains 10-15% trienleium aluminate, this is minimized to 2% in seawater cement.

Printers' Lead Putties, also termed Hard Putty and Carringe Putty, will vary in lead content from almost straight lead to approximately 75 per cent and 50 per cent; the admixtures being whiting and/or sahen. Typifying the first two, are the formulas below of hard putters actually used in railrond shops:

Dry White Lead	90 lb.	50	lb.
White Lead in Oil		20	lh.
Whiting		25	lb.
Silex	6 lb.		
Boiled Linseed Oil	3'in ga	l	
Gold Size Japan	3'ig ga		gal.
Rubbing Varnish	1 1/4 gal	1. 3/4	gal.

These mixtures are allowed to stand 72 hours to thoroughly wet down and swent, and then kneaded up into putty. The silex used is the live quartz silica namb adopted for the making of paste wood fillers. The pigmentation of a representative painters' hard putty with lower lead content would be 50% dry white lead, 35% whiting and 15% silien.

A non-shrinkable type of putty containing about 20% of lead in the pigment is this:

Whiting	125	lb.
White Lead, Dry	371/2	lb.
Silien	121/6	lb.
Raw Linseed Oil	31/2	gal.
Flour Paste	101/4	

The flour paste is 2 lb, of wheat flour beaten up in about 1 quart of cold water and then poured into 3 quarts of boiling water, and boiled 5 minutes. Yield 101/4 lb, net.

The foregoing non shrinkable putty is very similar to what used to be known as Swedish putty, purported to be so excellent for wood, iron, or stone. Another type of Swedish Putty without lead, is the following:

Rye Flour	2	lb.
Cold Water		gal.
Beat together, then pour	into	
Boiling Water	1	gal.
Cook 5 minutes, let cool,	then	stir

Whiting	20	lb.
Whiting Gold Size Japan Raw Linseed Oil Grind in a paint mill	50 2 1	lb. gal. gal.

Combine the two parts in a pony chaser, and thicken with more whiting to the required plasticity for knifing. This batch produces 100 lb. net.

Metal Furniture Baking Putty
Bolted Whiting 5 lb.
mixed with
Boiled Linseed Oil 1 pt.

then
Flour Paste 1 pt.

Mix all very thoroughly. The flour paste is as given for non-shrinkable putty. In all cases of preparing flour pastes, the flour and cold water should be beaten until entirely free from lumpiness; and during the subsequent cooking, should be continually stirred.

Stopping Putty is a dry mixture of 2 lb, of "Alabastine," 1 lb, of wheat flour, and 1 lb, of Portland Cement. When ready to use, 1 pound of this mixed powder should be thoroughly worked up to a stiff putty with 8 fluid ounces (½ pint) of cold water. This putty sticks to stone, wood, brick, etc.; used for filling knot holes, cracks, etc. Keep the dry powder in an nir-tight jar.

Gesso Duro is Italian hard plaster used in making bas-relief easts. When dried, it becomes very hard and durable.

This product, per formula, below, remains soft and manipulable for quite a period of time, using a small trowel, spatula or by forming with the hands:

LePage's Fish Glue gal. Water, to reduce it Oil of Lavender gal. 6 fl. oz. Raw Linseed Oil gal. 1 Bolted Danish Whiting ĺb. 50 Rubbing Varnish Bolted Danish Whiting gal. lb. 1 20 (colors in oil may be added, if shading is desired)

Plastic Wood Dough

*Gum Solution	1 gal.
Glycerin	3 pt.
Butyl Alcohol	3 pt.
Whiting	8 lb.
Wood Flour	24 lb.
Dope (Solution)	8 gal.
*The ((mun') solution i	a 16 nounds o

*The "gum" solution is 16 pounds of gum rosin (WW Rosin) cold-cut (dissolved) in 1 gallon of methyl acetone; the "dope" is another cold-cut solution, basis of 1 pound of "movie" film scrap to each gallon of methyl acetone. The picture film scrap should be desilvered by washing in hot water to remove its gelatin coating and then laid out in the sun and air to dry; but preferably it is obtainable cleaned and ready for cutting.

Onyx Cement

The above wood dough product is a soft workable putty easily applied to all kinds of depressions to be surfaced up. The work or job should not be left in too-rough state because the putty dries and hardens very rapidly; the ultimate sanding down later is a rather tough job unless the puttying had been reasonably smoothly applied.

There is one putty specially used in fair quantity, which is very little known in regular paint circles. This is termed Onyx Cement because its specific utility is for bonding slabs of onyx, marble, glass, and their imitations, to the walls in public buildings. It is necessarily of rather firm plasticity because of the weight it must partially support. Uniform handfuls of the putty are attached to the wall foundation at intervals about 18 to 24 inches apart; the slabs mentioned are then stood upright on their base, and then pressed back steadily and firmly into the mounds of putty. Suction, and the adhesive strength of the putty, securely hold the murble and glass permanently in place. The same material, plain or colored, is embedded in the joints between the slabs. The composition of this putty follows:

Domestic Whiting, 350 Mesh	100	lb.
Domestic Whiting,	100	10.
200 Mesh	100	lb.
"Super-Sublimed"	White	
Lead	40	lb.
White Oil Drier		gal.
Bodied Linseed Oil		gal.
Boiled Linseed Oil	21/2	gal.

For certain work a Black Onyx Cement is used. This is produced on a bituminous base.

Another specialty probably even less known than the onyx putties . . . in paint circles, is a Black Packing Compound required by makers of corrugated iron culverts. These culverts are sturdy Armeo-iron corrugated pipe, galvanized, in sizes from 12 to 84 inches diameter. They are the aqueducts for streams crossing the highways and for surface-sewers under driveways in rural districts, etc. There is first applied hot a thoroughly-

tested bituminous mastic pavement along the line of flow where erosion is greatest ... approximately the lower one quarter or one-third of the inside circumference. This coating practically fills the valleys of the corrugations and to the extent of building up a thickness of perhaps 4-inch over the rises.

For this purpose the culvert manufacturer supplies a plastic for cold application. The composition is 3 parts by weight of sawdust and 1 part ashestos fiber, thoroughly churned together with enough coal tar solution to form a putty that may be applied by hand to the abraded spots in the paved section of the culvert.

The last unusual specialty to be mentioned is Sheet Metal Deadener. Two eastern manufacturers have been supplying during the past three or four years a plastic compound developed for sounddeadening sheet metal equipment, principally metal furniture and automobile parts. This became most essential with the advent of the closed body, to ehminate rumble and vibratory noises, and especially the "tinny" sound caused by closing the doors. It is a standard application on Ford, Auburn, Stutz, Marmon, Duesenberg, and Nash cars; and probably on many others. The material might be described as a very soft bituminous plastic apparently containing

fine asbestos fiber or other filler; it sur faces dust free very quickly, has excel lent adhesion and undoubtedly maintains flexibility indefinitely. As general practice, it is applied onto the inner surfaces of the auto body and doors, or other ob ject, to a thickness of approximately 1/4inch, using a trowel, broad knife, or spatula. This sets in less than 30 minutes, but soft; is firm in 11/2 hours and still somewhat soft, is solid in 4 hours but not hard: and shrinks down somewhat in solidifying. For large production as by body builders and in the auto plants, the material has sufficient "slip" so it can be sprayed with special equipment.

High grade cork paint films insulate surfaces against heat, cold, and moisture, also deaden sound and soften the effect of shocks and blows, rendering them valuable for use on automobiles, railroad cars, and aeroplanes. In the automobile industry they are employed to advantage on the lower sides of the engine bonnets and mud guards. Applied to the bonnets, they protect the outside lacquer films against the radiating heat of the motor; while the cork paint films on the lower sides of the mud guards protect the latter against the impact of stones, sand, etc. Applied to the surfaces of aeroplane cabins, they form a rather effectual insulation against the

noise of the motors.

COATINGS, PROTECTIVE AND DECORATIVE

Marine Paints

Marine paints differ from house paints chiefly in that harder pigments are required. This means that such pigments as zinc oxide and iron oxide are used more extensively in marine paints than in house paints. Since steel vessels have largely replaced wooden vessels in seagoing traffic, the formulas shown herein are for the preservation and beautifica-tion of steel rather than wood. On steel the priming coat of paint—that is, the paint applied first on the metal—is of more importance than the priming coat on wood. The service to which marine paints are exposed is much more severe than that to which house paints are exposed. To meet this condition the various parts of the vessel must be considered separately. The paints suitable for the parts seen from the outside when the vessel is affoat are quite different from the paints suitable for underwater portions of the vessel. The paints suitable for inboard bulkheads are quite different from those suitable for inner bottoms or bilges, etc.

An excellent priming paint for steel surfaces to be exposed to the atmospheric elements is made from the following formula which produces one gallon and spreads approximately 650 sq. ft. per gallon:

20 lb.
5 pints
2 gills
2 gills

Paint from the above formula should be used within a month after it is mixed. If allowed to stand in closed (or open) containers for an appreciably longer period, the pigment settles hard and cannot be again strired to proper consistency for painting. By using very finely ground red lend pigment which contains 99 per cent true red lead, it is possible to successfully store the paint through periods of approximately one year. However, if the paint is to be stored during such period, or longer, formulas such as the following should be used:

OHOWING SHOULD BE WIND	Y .			
Red Lead, Dry	* 1	lb.	11	oz.
Zinc Oxide, Dry			13	oz.
Venetian Red, Dry	4	lb.	2	oz.

Spar Varnish	2 lb.
Raw Linseed Oil	2 lb. 7 oz.
Petroleum Spirits	9 oz.
Paint Drier	14 oz.
Aluminum Stearate	1 oz.
Films from paints	of the above for-
mulas interfere with	the adhesion of
shipbottom paints, so t	hese paints should
not be used on the o	utside underwater
portion of the hull.	If it is desired to
prevent corrosion on the	hat portion of the

10 07

Magnesium Silicate, Dry

Metallic Brown, in Oil	7.5 lb.
Raw Linseed Oil	2.3 lb.
Spar Varnish	.3 lb.
Gasolino	.6 lb.
or	

vessel during construction, a weaker film

paint should be used, such as:

Gasolino		.6	Ib.
	or		
Metallie Brown,	Dry	4.0	lb.
Spar Varnish	•	4.4	lb.
Paint Drier		2.5	lb.

The above two formulas are also suitable for a paint to be used on freshly pickled steel to protect it during fabrication; that is, as shop coat or field coat paints.

Aluminum paint may be used in lieu of red lead paint, for priming steel, but should not be used on underwater portions of the vessel. Its bright luster aids inspection of the interior of vessels under construction, but in warm, humid clumates it does not prevent rust as does red lead paint. The formula is:

Aluminum Powder 2 lb.
Aluminum Mixing Varnish 1 gal.
Note: This paint should be used within a few hours after mixing.

While priming paints will give fair protection when used alone, they are designed to be covered with at least two coats of finishing paint. Unlike house paints, there is no advantage in using a different formula for the first and the second coat of marine finishing paint. Following are formulas for ten gallons of finishing paints—on surfaces not to be exposed underwater:

Outside White Paint

Titanox	B, in	Oil	85	lb.
Zine Ox			36	lb.

Ultramarine Blue, in Oil	.5	oz.	Paint Drier	4	
Raw Linseed Oil	30		Ultramarine Blue, in Oil	.5	OZ.
Petroleum Spirits		lb.			
Paint Drie	8 1	lb.		_	
or			Inside White Enam	el	
White Lead, in Oil	53 1	ъ.	Titanox B, Dry	25	lh.
Zinc Oxide, in Oil		b.	Zine Oxide, Dry	25	lb.
Raw Linseed Oil		b.	Damar Varmsh	68	lb.
Petroleum Spirits		b.	Pine Oil	ű,	lb.
Paint Drier		b.	Ultramarine Blue, in Oil	.3	OZ.
Ultramarine Blue, in Oil		oz.	To this white enamel may	y be s	hobba
Citiana in a series		1	color pigments, ground in oi	l or i	n var
O till Divi Dele			nish, to produce desired shade	's. 153	Rad.
Outside Black Pain		. 1	ing additional pine oil just b	efore r	ippiy-
Lampblack, in Oil		b.	ing, the enamel is made to	o bru	sh on
Raw Linseed Oil		b.	much easier. An cuamel will	not a	dhere
Paint Drier	14	b.	well over an enamel or gloss	y nnisi	11. 11
or		.	two coats are to be applied, t	ne nrs	I cont
Lampblack, Dry		lb.	should be a flat paint.		
Spar Varnish		b.			
Petroleum Spirits		lb.	O Aut 1 Durit Dain		
Paint Drier	18	lb.	Outside Buff Pain	125	њ.
			White Lead, in Oil	14	lb.
Inside White Paint	t .	. 1	Yellow Ochre, in Oil	5	lb.
Titanox B, in Oil	76 51 45 8 20	lb.	Venetian Red, in Oil	27	lb.
Zinc Oxide, in Oil	51	lb.	Raw Linseed Oil	7	ib.
Raw Linseed Oil	4.5	lb.	Petroleum Spirits	4	lb.
Damar Varnish	. 8	lb.	Paint Drier	-	10.
Petroleum Spirits	20	lb.			
Point Drier	-1	11).	Inside Semi flat Light Gro	on Pa	int
Ultramarine Blue, in Oil	.5 €	oz.		65	lb.
or			Titanox B, Dry	30	lb.
White Lead, in Oil		lb.	Zine Oxide, Dry		07.
Zinc Oxide, in Oil		lb.	Chrome Green Oxide, in O	39	lb.
Raw Linseed Oil		lb.	Damar Varnish	20	lb.
Petroleum Spirits		lb.	Petroleum Spirits	20	1.5.
Paint Drier		lb.			
Ultramarine Blue, in Oil	.5	oz.	Inside French Gray E	namel	
			Titanox B, in Oil	72	lb.
Light Gray Paint	;		Lampblack, in Oil		lb.
Titanox B, in Oil		lb.	Chrome Yellow, in Oil	1	1b.
Zinc Oxide, in Oil		lb.	Spar Varnish	30	lb.
Lampblack, in Oil	1 3/4	lb.	Damar Varnish	29	lb.
Ultramarine Blue, in Oil	3/4	lb.	Pine Oil	6	lb.
Raw Linseed Oil		lb.			
Petroleum Spirits	1.5		Piping, ducts, gas cyl	indora	etc.
Paint Drier	8	lb.	aboard vessels are usually	marke	l with
			colors to indicate the purpe	se ser	ved or
Outside Green Pair	nt		the contents. Formulas for	such	paint
		lb.	are:		
Chrome Green, Dry		lb.	Red Paint		
Zinc Oxide, Dry	3.6		1		7 lb.
Chrome Yellow, in Oil Yellow Ochre, Dry Lampblack, in Oil	7.5		Toluidine, Dry		3 lb.
Lampblack, in Oil		lb.	Spar Varnish	1.	
Spar Varnish	35	lb.			
Petroleum Spirits		lb.	Blue Paint		
Paint Drier	4	lb.	i .	10	6 lb.
I dint Ditt.			White Lead, in Oil		6 lb.
- 13 TH (1771 17 Th	aint		Ultramarine Blue, in Oil		2 lb.
Inside Flat White P		,,	Raw Linseed 31 Petroleum Spirits		8 lb.
Zinc Oxide, in Oil	157	lb.	Paint Drier		4 lb.
Petroleum Spirits	23	lb.	1 Bills Direct		

26 THE CHEMICAL FORMULARY			
Green Pair	nt	Green Pain	t
Chrome Green, in Oil	97 lb.	Chrome Green, Dry	30 lb.
Raw Linseed Oil	21 lb.	Lampblack, in Oil	2 lb.
Petroleum Spirits	9 lb.	Interior Varnish	• 60 lb.
Paint Drier	5 lb.	Paint Drier	10 lb.
Tant Driei	- 0 10.	Tallit Dilei	- 10 10.
Black Pain		Black Paint	;
Lampblack, in Oil	70 lb.	Drop Black, Dry	38 lb.
Petroleum Spirits	9 lb.	Interior Varnish	48 lb.
Paint Drier	10 lb.	Paint Drier	16 lb.
Brown Paint	- :	m	
Metallic Brown, in Oil	100 lb.	The waterline area on t	he outside of
Raw Linseed Oil	27 lb.	the hull is generally reg	arded as the
Petroleum Spirits	8 lb.	most difficult part of the	ressel to keep
Paint Drier	3 lb.	properly painted. This is	because it is
I am Dilei	3 10.	subjected to both atmospher	ic and under-
77.11 75.1		water exposure, and paints	suited to the
Yellow Paint		one exposure are not suited	to the other.
Chrome Yellow, in Oil	116 lb.	A high grade varnish paint	applied over
Raw Linseed Oil	20 lb.	red lead primer gives as goo	od service on
Petroleum Spirits	9 lb.	this area as has been obtain	ed. Typical
Paint Drier	3 lb.	of waterline paints are:	
		Red Paint	
The above red and green	paints are	Venetian Red, Dry	29 lb.
itable for the stands on w	hich running	Spar Varnish	42 lb.
thts are mounted, red mark		Petroleum Spirits	7 lb.
de and green the starboard		Paint Drier	18 lb.
Single shell smoke stacks	become too		
t for any of the above p	aints. Such	Light Gray Pain	t
rfaces should be painted	with special	Zinc Oxide, Dry	30 lb.
ints, the following for	nuias being	Lampblack, in Oil	8 lb.
pical:		Ultramarine Blue, in Oil	12 lb.
Light Gray Pain	t i	Spar Varnish	31 lb.
White Lead, Dry	48 lb.	Petroleum Spirits	17 lb.
Zinc Oxide, Dry	19 lb.	Paint Drier	18 lb.
Litharge, Dry	3.5 lb.		
Lampblack, in Oil	.5 lb.	Black Paint	
Ultramarine Blue, in Oil	.5 lb.		20 lb.
Damar Varnish	20 lb.	Drop Black, in Oil Zinc Oxide, Dry	19 lb.
Kerosene	33 lb.	Spar Varnish	20 lb.
Paint Drier	6 lb.	Petroleum Spirits	20 lb.
or or		Paint Drier	17 lb.
Titanox B, Dry	60 lb.	or	11 10.
Interior Varnish	52 lb.	Lampblack, Dry	8 lb.
Lampblack, in Oil	2 lb.	Zine Oxide, in Oil	20 lb.
Petroleum Spirits	9 lb.	Spar Varnish	38 lb.
		Petroleum Spirits	7 lb.
Red Paint		Paint Drier	18 lb.
Indian Red, Dry	40 lb.		
Interior Varnish	55 lb.	Shipbottom paints are use	to prevent
Paint Drior	15 lb	Surpostion burnes are nace	a so biesent

Shipbottom paints are used to prevent rust and to prevent the attachment of marine fouling on the bottoms of vessels. The "anti-corrosive" paint is to prevent rust and is applied next to the steel. The "anti-fouling" paint is to prevent the attachment of barnacles, algae, and other forms of fouling. It contains meterial forms of fouling. It contains material toxic to marine organisms, and is applied over the anti-corrosive paint. Both paints should be quick drying paints. Each of the two paints is so dependent on the

Buff Paint

lb.

lb. 55 55 18 13 lb. lb.

lb.

12 lb. .5 lb. 3 lb. 5 lb.

23 15

Paint Drier

Silica

White Lead, Dry White Lead, in Oil

Yellow Ochre, in Oil

Litharge, Dry
Venetian Red, in Oil
Boiled Linseed Oil
Petroleum Spirits

		TO MILE IN	OMMITTI		
other that the two formulas	are shown	R	lack Deck Pai	int	
together. The anti-corrosive p		I samueli i	, Dry ish Spirits		4 lb.
set should not be used with		Lampoiack	, 17ry		
fouling paint of another set.	The follow-	Spar Varn	ish .	4	4 lb.
ing formulas are typical:	The Tollow.	Petroleum	Spirits		5 lb.
		Paint Drie	r	1	8 lb.
Anti-corrosive Pain			ray Deck Pai	int	
Gum Shellac	8 lb.				11
Denatured Alcohol	54 lb.	Zine Oxide	, Dry , Dry	33	lb.
Zinc Oxide, Dry	29 lb.	Lampblack	, Dry	6_	lb.
Zinc Dust	11 lb.	[Ultramarin	e Blue, in Oil	1,	4 lb.
Denatured Alcohol Zinc Oxide, Dry Zinc Dust Pine Oil	5 lb.	Spar Varni	18h	7.1	
A 41 A II - D-1 A		Paint Drie	г	1	Ъ.
Anti-fouling Paint		_		•	
Gum Shellac	14 lb.		led Deck Pair		
Denatured Alcohol Zinc Oxide, Dry Indian Red (Iron Oxide)	45 lb.	Red Lead,	Dry	10) lb.
Zinc Oxide, Dry	14 lb,	Indian Red	(Iron Oxide),	, Dry 23	5 lb.
Indian Red (Iron Oxide)	15 lb.	Alummum	Stearate		2 lb.
Mercuric Oxide	8 lb.	Lamoblack.	in Oil		2 lb.
Pine Oil	9 lb.	Spar Varni	alı	4	4 lb.
Anti-fouling Paint, shown	abovo is	Paint Drier	Stearate in Oil sh	i	3 lb.
used with the Anti-corresive P	aint shown		e this paint		
above.		lead it can l	be unplied di	roetly o	in the
		steel deck; th	at is no red l	end pri	mer is
Anti-corrosive Paint		necessary.			
Zinc Oxide, Dry Venetian Red, Dry Silica	19 lb.				
Venetian Red. Dry	9 lb.	Black	Anchor Chain	Paint	
Silica	9 lb.		ALL HOL CHAIN		7 1b.
Rogin (WW Grade)	15 lb.	Gilsonite			
Solvent Norththa	38 lb.	Rosm) ()Z.
Manganaga Lindonta	13 lb	Petroleum l	Residuum phtha	2	l Ib.
Silica Rosin (WW Grade) Solvent Naphtha Manganese Linolcate Coal Tar	5 lb	Solvent Naj	phtha	4	7 lb.
Anti-fouling Paint	04.11	Green	Anchor Chain	Paint	
Zinc Oxide, Dry	24 lb.	Chrome Gre	en, in Oıl Dry	10	lb.
Asbestine, Dry	7 lb.	Red Lead,	Dry	10	lb.
Silica	8 lb.	Aluminum	Powder	5	lb.
Cuprous Oxide	15 lb.	Asohaltum	Varnish	4	gal.
Mercuric Oxide	4 lb.	Roiled Lans	eed Oil	2	gal.
Rogin (WW Grade)	25 lb.	Spar Varni	sh	2	gal.
Solvent Numbths	34 lb.	Potroloum	Smirita	5	mil
Diag Oil	4 lb	Deint Dage	- milita	1/-	gal.
Coal Tar	6 lb.	Taint Drie	en, in Oıl Dry Powder Varnish seed Oıl sh Spirits	- 72	gai.
Anti-fouring Faint Zinc Oxide, Dry Asbestine, Dry Silica Cuprous Oxide Mercuric Oxide Rosin (WW Grade) Solvent Naphtha Pine Oil Coal Tar			e used to brig		
		work on mar	ing regards	The fee	lowing
red lead paint and finished	with two	work on man	for vessels.	The 10	HOWING
coats of one of the following de	ck paints:				
		Orange	Red Shellac 27 lb. 48 lb. 17 lb.	Gre	en
		(clear)	Shellne	Shel	lac
		21.16	27 lb	97	115
Gum Shellac		61 III.	40 11.	Shel 27 53	11.
Wood or Denatured Alcoh	01	5.5 ID.	17 11.		11).
Venetian Red (Iron Oxide	')		17 10.	15	11
Chrome Green, Dry			-	217	
Drop Black, Dry		-		15	10.

20 lb. 15 lb. 5 lb. 2½ lb. 39 lb. 5½ lb.

Bilge and Tank Paint			Black Acid Resisting	Paint 20
Black Flexible Paint			Paving Asphalt	15
etroleum Residuum tosin	7	lb. lb.	Lampblack, Dry Beeswax	5 21/2
Petroleum Spirits		lb. lb.	Petroleum Spirits Paint Drier	39 51/2

Bituminous Enan	nel	Cobalt Paint Dr	ion
Petroleum Residuum	80 lb.	Cobalt Acetate	
Paving Asphalt	10 lb.	Pusin Fatan Com	5 lb.
Asbestos Fiber	5 lb.	Pow Lingued Od	15 lb.
		Rosin Ester Gum Raw Linseed Oil Petroleum Spirits	19 lb.
Note: This product must	be neated for	retroieum spirits	39 lb.
application.			
Potable Water Tank	Doint	Asphaltum Varn	ish
Mark Water Tank	raint	Paving Asphalt 3	5 lb.
Metallic Brown, Dry	40 lb.	Paving Asphalt Manganese Resinate Litharge Raw Linseed Oil Petroleum Spirits 3	7 lb.
Indian Red, Dry	15 lb.	Litharge	1 lb.
Zinc Oxide, Dry	8 lb.	Raw Linseed Oil	5 lb. 5 oz.
Manhami Mannish	8 lb.	Petroleum Spirits 3	9 lb.
Detailmen Cairite	94 ID.		
Point Drive	3 ID.		
Metallic Brown, Dry Indian Red, Dry Zinc Oxide, Dry Silica *Amberol Varnish Petroleum Spirits Paint Drier	3 ID.	Damar Varnish	
		Batavia Damar Gum Turpentine	47 lb.
Raw Tung Oil	35 B	Turpentine	22 lb.
Amberol Gum No 226 Raw Tung Oil Petroleum Spirits Cobalt Drier	39 lb.	Petroleum Spirits	21 lb.
Cobalt Drier	₹4 lb.		
Tol 1 m 1 75 1		Copper Paint for Wood	Bottoma
Black Tank Pair	nt	Gum Shellac Denatured Alcohol Zinc Oxide, Dry Indian Red, Dry Cuprous Oxide Pine Oil	16 lb
Petroleum Residuum	12½ lb.	Denstured Alcohol	50 lb.
Litharge	13/4 lb.	Zane Oxide Dry	161/ lb
Red Lead	1¼ lb.	Indian Red Dry	1614 lb
Rosin (D Grade)	1/4 lb.	Cuprous Oxide	8 lb
Lampblack, Dry	5 1/2 16.	Pine Oil	9 lb
Petroleum Residuum Litharge Red Lead Rosin (D Grade) Lampblack, Dry Boiled Linseed Oil Spar Varnish Damar Varnish Petroleum Spirits	12 lb.		0 10.
Spar Varnish	14 lb.	Anti Pouling Waterla	- D. t
Daniar varnish	4 lh.	Anti-Fouling Waterlin	
Petroleum Spirits	32 ½ 1b.	Gum Shelhac Denatured Alcohol Pine Oil Crude Rubber Gasoline Zine Oxide, Dry Lampblack, Dry Moreure Oxide Turpentine	13 lb.
		Denatured Alcohol	5 gal.
Brown Tank Pai	nt	Courts Bulker	3 gal.
Metallic Brown, Dry	40 lb.	Canalina	1 OZ.
Litharge	2 lb.	Zina Ovula Dan	z gms
Zinc Oxide, Dry	16 lb.	Lamphlack Dry	0 10,
Zinc Chromate, Dry	2 lb.	Moreura Ovida	4 10,
Damar Varnish	46 lb.	Turpentine	2 lb.
Interior Varnish	11 lb.		2 IU.
Metallic Brown, Dry Litharge Zinc Oxide, Dry Zinc Chronate, Dry Damar Varnish Interior Varnish Paint Drier	15 lb.	White Water Pai	_4
		Zina Ovido Day	24 lb.
Primer for Bituminous		Zine Oxide, Dry Whiting, Dry	40 11.
Trinidad Asplult	53 lb.	Plaster Pura	9.1 lb
Petroleum Spirits	6% gal.	Pulverized (Hide) Glue	4 lb.
		Plaster Paris Pulverized (Hide) Glue Ultramarine Blue, Dry Note: Mix 8 lb, of the al	1 07
Bituminous Enam	el	Note: Mix 8 lb. of the al	hava mirtura
Paving Asphalt Trinidad Asphalt Rock Asphalt Rosin (Dark Grade) Portland Cement Slacked Lime	52 lb.	in one gallon of water.	DOTE MIXIME
Trinidad Aspbalt	15 lb.	in one gamon of water.	
Rock Asphalt	15 lb.	-	
Rosin (Dark Grade)	1 lb.	White Enamel	
Portland Cement	17 lb.	Titanox B, Dry	72 lb.
Slacked Lime	21/2 lb.	Spar Varnish	28 lb.
Note: This product must l	oe heated be-	Spar Varnish Damar Varnish	29 lb.
fore applying.		I Inc On	0 10.
		Ultramarine Blue, in Oil	1 oz.
Paint Driers			
Manganese Resinate	10 lb. 10 lb. 2 lb. 8 lb.	Gray Enamel	
Damar Gum	10 lb.	Titanox B, Dry	60 lb.
Litharge	2 lb.	Lampblack, in Oil	2 lb.
Raw Linseed Oil	8 lb.	Interior Varnish	53 lb.
Littiarge Raw Linseed Oil Petroleum Spirits	49 lb.	Titanox B, Dry Lampblack, in Oil Interior Varnish Petroleum Spirits	9 lb.
		-	

COATING	S, PROTECT
Red Ename	
Indian Red, Dry	40 lb.
Interior Varnish	
Paint Drier	55 lb.
	9 lb.
Petroleum Spirits	6 lb.
Outside White I	`aint
Zinc Oxide, in Oil	50 lb.
Basic Sulphate White Le	and
in Oil	50 lb.
Blanc Fixe, in Oil	12 lb.
Asbestine, in Oil	6 lb.
Raw Linseed Oil	4 gal.
Petroleum Spirits	¼ gal.
Paint Drier	1/2 gal.
Ultramarine Blue, in Oil	1 oz.
Red Lead Pan	ıt
Red Lead, Dry	85 lb,
Silica	40 lb.
Raw Linseed Oil	61/4 gal.
Petroleum Spirits	5, gal.
Paint Drier	⅓ gal.
Light Gray Par	nt
Zinc Oxide, Dry	34 lb.
Blanc Fixe, Dry	34 lb.
Graphite, Dry	2 lb.
Lamphlack in Oil	1 oz.
Lampblack, in Oil Ultramarine Blue, in Oil	
Raw Linseed Oil	6% gal.
Petroleum Spirits	1 gal.
Paint Drier	3/4 gal.
The formulas shown rec pigments in oil be stiff pas centages of raw linseed o within the limits shown:	pure that the tes. The per all present are
	in l'aste
White I and (Carboneta)	8 to 10
White Lead (Carbonate)	8 to 10
White Lead (Sulphate)	
Zinc Oxide	8 to 18
Titantium Pigment B	15
Chrome Green	33 to 35
Chrome Oxide, Green Chrome Yellow	29 to 31
Chrome Yellow	24 to 26
Metallic Brown	22 to 24
Lampblack	65 to 80
Raw Sienna	45 to 55
Burnt Sienna	40 to 50
Raw Umber	35 to 45
Burnt Umber	30 to 50
	30 to 40
Yellow Ochre	20 to 30
Magnesium Silicate	20 to 30 20 to 25

20 to 25

15 lb. 25 lb.

35 lb.

10 lb.

Venetian Red

Carbon Black

Barytes Boiled Linseed Oil

Kaolin

Black Marine Paint

The second secon	
Red Paint	
Indian Red	5 lb.
Barytes	1 lb.
Whiting	1 lb.
Linseed Oil	2 lb.
Japan Drier	6 oz.
Mixing Varnish	5 lb.
Surfacer	
Varnish	1 gal.
Brown Japan	1 gal.
Silex (Fine)	8 lb.
Ship Bottom Paints	
1. For Wood Bettoms	
In any formulation, the obje	ct should

In any formation, the object some be, first, to produce a mixture which will best serve the purpose and, second, to obtain the mixture at the lowest cost. The work requires a knowledge of a wide range of materials, their chemical and physical properties, and their cost. It also requires a knowledge of paint manufacturing operations, especially those to which the equipment on hand is adapted. Formulating is not an exact science any more than is the prescribing of medicine by the physician. One important difference between the physician writing a prescribing a paint formula is that the latter is also thinking about the cost.

The requirements of a paint for wood bottoms are comparatively simple and casy to meet. The corrosion problem does not enter, and consideration of a possible chemical or physical conflict with a prining paint does not enter. The object is to produce a paint, the film of which will brush (or sprny) on casily, will dry quickly, will be resistant to water crossion and yet sufficiently softened by the water to permit the toxic elements to go into solution. There are several ways of approaching the problem which can best be illustrated by used formulas.

Formula No. 1

1 01111 445 1	10. 1	
Iron Oxide	18	lb.
Silica	5	lb.
Copper Cyanide	13.5	lb.
Spar Varnish	7.25	gal.
Pine Tar Oil	.625	gal.
Paint Drier	.23	gal.
"Tar Acid Oil"	.30	gal.
Moneral Spirits	.25	gal.

(Comment: The above formula will doubtless "dry" in about four hours because the spar varnish, which usually requires about twelve hours to dry, has been overloaded with the added driers. The dried film will be glossy and apparently hard, but it will probably not dry hard because of the excessive pine tar oil. The toxicant, copper cyanide, is regarded as only fairly toxic. This fact, together with the fact that a spar varnish film usually disintegrates under sea water and fouls readily, suggests that the film will not prevent barnacle fouling for a longer period than two or three

Following is a formula which has given very good service:

No. 2		
Blanc Fixe	40	lb.
Mercuric Oxide	5	lb.
Paris Green	7.5	lb.
Gum Shellac	20	lb.
Denatured Alcohol	5.9	gal.
Pine Oil	2.5	gal.

(Comment: The above formula is typical of shellac type paints. This paint will be effective about six months on a wooden bettom. It probably will not stand long storage satisfactorily, the nature of the pigment being such as to suggest a very hard sediment forming).

The U.S. Navy used a formula similar to the above.

No. 3	
Zine Oxide	165 lb.
Indian Red	165 lb.
Cuprous Oxide	75 lb.
Gum Shellac	162 lb.
Alcohol	500 lb.
Pine Oil	90 lb.

2. For steel bottoms.

In successfully formulating paints for steel bottoms the maximum ingenuity of the paint technologist is required. There are wide variations of opinions among men engaged in this work and each opinion is based, more or less, on experience in research. In designing paints for exposure to atmospheric elements there are certain fairly well established rules as to pigment vehicle ratios by weight and by volume. For an oil paint for outdoor exposure, the pigment should be about 60 per cent by weight, and about 29.5 per cent by volume, of the paint. No such rules have been, or can be, established for ship bottom paints. Such ratios vary with each change in the vehicle, and there are an almost infinite number of such changes that can be made. The setting of high and low limits for the variants is apparently useless.

Before considering the varnish type of paints, which general type constitute the bulk of ship bottom paints used in America, the hot plastic paints, such as are used extensively in European countries, will be considered. Following are formu-las used about twelve years ago by one of the European Navies.

Anti-corrosive Paint.

Rosin	26.5	lb.
Benzol	26.5	lb.
Ozokerite	5	lb.
Iron Oxide	42	lb.

Anti-fouling Paint

Rosin	38.6	lb.
Stearin	14.7	lb.
Benzol	12.8	lb.
White Lead	7.4	lb.
Verdigris	9.6	lb.
Arsenie	13.2	lb.
Mercuric Oxide	3.7	lb.

To illustrate the varnish type ship bottom paints, two sets of paints used by the United States Navy are shown.

Anti-corrosive Paint

Formula No. 1

One Gallor	n Formula	
Zinc Oxide	3.05	lb.
Zine Dust	1.1	lb.
Gum Shellac	.425	lb.

Gum Shellac			.425	lb.
Yacca Gum			.44	lb.
Alcohol			.8	gal.
Pine Oil			.067	
	NT.	0		•

rine On	.067 gal.
No. 2	-
Coal Tar	47.5 lb.
Rosin	145 lb.
Coal Tar Naphtha	380 lb.
Magnesum Linoleato	129 lb.
Venetian Red	93 lb.
Zinc Oxide	186 lb.
Silica	93 lb,
Beeswax	3.3 lb.

Anti-Fouling Paint Formula No. 1

One Gallen Formul

One Ganon Form	iuin	
Zinc Oxide	1.65	lb.
Indian Red	1.65	lb.
Mercuric Oxide	.75	lb.
Gum Shellac	.815	lb.
Yacca Gum	.89	lb.
Alcohol	.76	gal.
Pine Oil	.125	gal.

No. 2		
Coal Tar	132.6	lb.
Rosin	202.0	lb.
Coal Tar Naphtha	228.0	lb.
Pine Oıl	74.0	lb.
Zinc Oxide	212.0	lb.
Silica	82.0	lb.
Asbestine	83.0	lb.

Cuprous Oxide Mercuric Oxide 112.0 lb. 45.0 lb.

Although commercially made phenolformaldehyde condensates have not proved satisfactory in undersea water exposure, there apparently is considerable merit to a varnish from such resin when the resin is made simultaneously with the varnish. These varnishes comprise the vehicle of the ship bottom paints and are made in reflux condensers. Typical of the process is the following:

Place 90 lb. of phenol, 108 lb. of 40c; solution of formaldelyde, 90 lb. of water and 54 lb. of lead acetate in a refux condenser and boil about 30 minutes. Add 720 lb. of rosm and continue heat until excessive foaming starts. Remove the reflux and continue heat until foaming ceases and at same time blow air through the mixture. Cool and add 108 gal. of coal tar naphtha.

The varnish is mixed with pigments to form anti-corrosive and anti-fouling paints.

Anti-Corrosion and . Fouling Paint Yacca Gum 1.6 lb. Alcohol 1.32 gal. Pine Oil 1.9 gills Petroleum Spirits gills 1.9 Zinc Oxide 1.2 Ϊb. 1.2 lb. Silica Blanc Fixe 12 lb. lb. Zinc Dust 0.3Paris Green 0.6 lb. 1.4 lb. Mercuric Oxide

Paints for Ship Bottoms Formula No. 1

2.5 parts of wood tar, 2.0 parts of oxide of iron, 1.0 part of turpentine result. 2.0 parts of lead acetate. Wood tar is preferable to coal tar, since the latter is not as resistant towards the corrosive action of sea water.

No. 2

1.0 parts of lead arsenate, 1.0 parts of Scheele's green (copper arsente), 8.0 parts of ochre, 5.0 parts of turpentine resin, 3.0 parts of coal tur, 2.0 parts of Bakelite, 5.0 parts of oil of turpentine and 5.0 parts of white spirit.

No. 3

The so-called "Lucchini Paint": 30.0 parts of galpot (white resin produced from fir), 20.0 parts of turpentine resin, 2.5 parts of mercury arsenate, 20.0 parts of red arsenic, 30.0 parts of wood tar, 5.0

parts of manganese dioxide and 15.0 parts of oil of turpentine.

No. 4

600 0 parts of asphaltum or pitch, 480.0 parts boiled linseed oil, 120.0 parts of graphite, 120.0 parts of arsenic copper oxide and 640.0 parts of coal tar oil.

No. 5

48.0 parts of coal tar, 383.0 parts of tar oil, 146.0 parts of turpentine resu, 130.0 parts of manganese linoleate, 3.3 parts of beessux, 93.0 parts of Venetan red, 93.0 parts of infusorial earth and 187 parts of zine oxide.

No. 6

133 parts of conl tar, 288 parts of tar oil spirits, 20 parts of turpentine resin, 74 parts linseed oil, 21 parts of zine oxide, 82 parts of infusorial earth, 83 parts of magnesium silicate, 112 parts oxide of copper and 115 parts mercury oxide.

Ship Bottom Paints

An anti-corresive paint is prepared from 145 parts of orticiea futty neids, 120 parts shellac, 390 parts nleohol, 80 parts pine oil, 5 parts drier, 170 parts of iron oxide, and 160 parts of zine oxide.

The anti-fouling composition given is, 115 parts of ontices fatty acids, 129 parts shelhar, 430 parts alcohol, 80 parts pine oil, 5 parts drier, 170 parts of iron oxide, 100 parts of zine existence, 100 parts of copper, and 40 parts of vellow oxide of mercury.

To prepare these paints the shellae is dissolved in some of the alcohol, the pine on added, and then the pigments ground in, using a ball mill. When perfectly smooth, the fatty acids, mixed with the pine oil and the remainder of the alcohol, are added. The same nethod is used in the case of the anti-fouling composition except that the yellow oxide of mercury is not ground in but only mixed.

After storage these products appear very thick, but spread easily under the brish to give a thick flexible film. A little diluent can be added if necessary.

The cobalt, lead and manganese salts of the fatty acids of oiticiae oil, which are in the nature of driers, are prepared by converting the acids into soluble soaps and precipitating these by the acetate of the appropriate metal. The precipitate is carefully washed and dried in a carefully regulated oven in a current of carbon dioxide to prevent oxidation.

Paints for Glazing and Coloring Ceramics
Pigments (glass-powder, colored with
metal oxides)

Thinner:
Linseed Oil 5 lb.
Wood Oil 3 lb.

Artificial Mother-of-Pearl British Patent 426,554

Dibasic lead phosphate (PbHPO₄) prepared by adding phosphoric acid to a warm solution of a lead salt, if used in the form of very fine crystals, produces glistening or iridescent effects. The salt may be obtained in a very fine state of division by precipitation in presence of a water-soluble organic compound, and preferably under slightly acid conditions. Thus 24 liters of a solution of lead mtrate (3.3 kg. dissolved in 10 liters of water) are mixed with 4.8 liters of distilled water and 24 liters of 95 per cent alcohol; 2.6 liters of phosphoric acid (12 kg, of concentrated acid plus 50 liters of 90 per cent alcohol) are added all at once. The use of this lead phosphate in materials which are to be submitted to treatment with formaldehyde (e.g., casein products) is advantageous in that the salt is not affected by formaldehyde.

Enamel Opacifier British Patent 427,850

A fine white powder is obtained by heating at 1000° for several hours an intimate mixture of titanium dioxide 33.1, antimony pentoxide 44.5, and zine oxide 22.4%.

Pearl or Fish Scale Essence German Patent 603,487

Formula No. 1

a. Scales of Uklei Fish

Treat a with b in a stirring-machine for 30 minutes, pour off the upper suspension, and repeat the same treatment of a two more times. Now the scales are free of fish-silver, the suspension containing about 1500-2500 g. of this substance.

No. 2

a. Herring Scales, Norwegian 100 kg.
b. { Ethyl Propionate 150 l. Nitrocellulose 1.5 kg.

As in No. 1. Yields in fish-silver are quantitative, viz. 700-1000 g. crude material.

No. 3

a. Astrachan Scales
b. { Ethyl Acetate }
l. Nitrocellulose
3 kg.

Work as in No. 1, stirring two times for 20 minutes. 1200-1600 g. crude fishsilver can be obtained by centrifugal separation of the suspension.

No. 4

a. Uklei Scales (from Lake

Scutari, Albania) 1 kg.
b. Ethyl Acetate 1.5 l.
c. \(\) \(\) Acetyl Cellulose 25 g.

Alcohol, a little to dissolve completely.

As in No. 1 in smaller proportions. Should yield 1.4 liter suspension with 0.4% dry fish-silver.

Protective Coating for Hydrofluoric

Beeswax 1 oz. Paraffin Wax 4 oz.

Electric Lamp Coating U. S. Patent 1,941,990

The lamps are coated with a paste of Kaolin 50 g. Guignet's Green 200 g. Cadmium Sulphide 50 g. Boric Acid 160 g. Sodium Silicate (d. 1.015) 1000 cc.

Coating Lacquer for Fabrics

Nitrocellulose Wet (5.6 sec.)	12 g.
Diamond 'K' Linseel Oil	8 g.
Crude Crepe Rubber (Light)	8 g.
Ethyl Acetato	10 g.
Butyl Acetato	10 g.
Aleohol	5 g.
Toluol	47 g.
Heet awale rubbes in birseel oil put	

Heat crude rubber in linseed oil until dissolved. Cool and dilute with part of toluol. Add to remainder of formula after nitrocellulose is dissolved.

Rubberized Cloth Varnish Formula No. 1

Shellac	5	oz.
Alcohol	95	0 z .
Gives high gloss.		

No. 2

Shellac Ammonia Water	(28%)	1 kg 1/2 kg 30 kg	١.
*******			•

Two coats of this must be applied to get good adhesion. The flush is semiglossy. These varnishes are applied by a velvet covered brush or roller.

Waterproofing Brick Walls

Walls can be waterproofed by applying a coat of solution made by dissolving 1% lb. of paraffin in each gal. of mineral spirits used as a solvent. Use steam to melt rather than a free flame.

Moisture proofing Compositions Canadian Patent 352,183

Moistureproofing compositions consist of (parts by weight): Formula No. 1, paraffin 85, refined caranuba wax 10, rubber 5; (No. 2) paraffin 65, rubber 5, candelila wax 30; (No. 3) paraffin 75, rubber 5, gum damar 20; (No. 4) paraffin 40, rubber 5, caranuba wax 40, ester gum 12; (No. 5) paraffin 60, rubber 5, caranuba wax 20, gum damar 15; and (No. 6) paraffin 55, rubber 4, candelila wax 25, hydrogenated castor oil 16 parts.

Jute Waterproofing French Patent 763,402

Asphalt	60 lb.
Bitumen	10 lb.
Coal Tar	5 lb.
Coal Tar Pitch	5 lb.
Linseed Oil, Boiled	2 lb.
Sand, Fine	15 lb.
Bordenux Resin	3 lb.

Straw Lacquer Waterproofing Italian Patent 267,765

Cellulose Nitrate	10	οz
Butyl Acetate		oz.
Benzol		OZ.
Butyl Alcohol		oz.
Paraffin Wax		υZ
Camphor Oil		OZ.
Butyl Ether	9	OZ.

Waterproofing Compound and Paint Vehicle

U. S. Patent 1,965,042

Three gallons of china wood oil is raised to a temperature of about 210°C, at this temperature 12 grams of manganese borate is added with rapid stirring. The temperature is maintained for a period not exceeding about fifteen minutes, but preferably from one to two minutes. In order to quickly cool the oil and also to partially dilute it, about 1 gallon of water white kerosene is added.

The temperature of the mass will thus be reduced to about 175° C. and when this temperature is attained 1½ pints of carbon tetrachloride is gradually added by introducing the same preferably near the bottom of the vessel. The rate of introduction of carbon tetrachloride is such that from 1–2 minutes are required for this step of the process. When the carbon tetrachloride has been introduced and the temperature has been reduced sufficiently, for example, to about 100° C, any desired quantity of diluent such as kerosene or solvent multiha is added.

This forms a solution of waterproofing material which when applied to stone, brick, masoury and the like penetrates the pores of the same and coats the surface of the material to which it is applied, efficiently protecting it from the clements such as rain, sea water, sail water air, heat and frost. The coating is not substantially acted upon by alkalies or neids and forms a colorless waterproofing material which remains effective for many years.

Waterproofing Composition Belgian Patent 400,446

The composition contains carbon tetrachloride or carbon disulphide 200 cc., paraffin 150 g., rubber 8 g., and naphthalene 50 g. per liter.

Waterproofing Composition U. B. Patent Serial Number 513,225

A waterproofing composition which comprises forming a mixture of from 285 to 290 parts of water, 12 to 16 parts of sodium sheate and 9 to 10 parts of oleic acid and then strring into this mixture approximately 300 parts of comminuted cumar resin (melting point about 230° to 245° F.) while maintaining the liquid at a temperature above 90° F. and not to exceed substantially 160° F.

Moisture and Greaseproof Coating

rormuta no. 1		
Gelatin	5.4	oz.
Bulfonated Oil	2.7	oz.
37% Formaldehyde Solution	1.4	oz.
Glycerin Monophthalate		
Ester	4.5	oz.

No. 2 In another specific formula, to each 100 oz. of a vehicle containing 10% alcohol add the following:

Gelatin 7.2 oz. Glycerin 3.6 oz. 58 07

37% Formaldehyde Solution 1.8 oz. Glycerin Monophthalate
Ester 3.6 oz.

No. 3

In a third specific example add to each 100 oz. of vehicle:

37% Formaldehyde Solution 1.4 oz. Glycerin Monophthalate, Elster 5.8 oz.

The two latter formulas, however, do not have the full effectiveness of the first in producing moisture-resistant and

greaseproof coatings.

Gelatin

In preparing the composition, when alcohol is employed in the vehicle, it is kept separate from the remaining constituents of the mixture until a late stage in the formation thereof. The gelatin is dissolved in a portion of the water, and, if desired, may be mildly acidulated, for example, with acetic acid. The flexibility-imparting agent, if any is used, is added to the aqueous solution of gelatin, suitably after admixture with or solution in a small amount of water, although this is not necessary. The formaldehyde solution is diluted with water. The di-luted formaldehyde solution is then added, or, in its place, suitable proportions of a solution of hexamethylenetetramine, or alum or the like may be employed. The alcohol is diluted, suitably with an equal amount of water, and then added to the mixture. The glycerin phthalate ester or other ester employed is then dissolved in part or all of the remaining quantity of water, neutralized, for example, with ammonium hydroxide, and incorporated in the mixture.

Waterproo:	f Finish	₩
For	f Finish rmula No. 1	No. 27
TOTHESIT	20 g.	20 g.
Methyl Abietate	12 g.	16 g.
Cumar V	12 g.	24 g.
Indian Red	25 g.	
Titanium Dioxide		40 g.

Waterproofing Fibrous Materials U. S. Patent 1,965,630

One thousand pounds of pulp fiber dry weight is mixed in an ordinary paper mill beater with about 20,000 pounds of water. To this is added about 300 pounds of alkaline filler such as calcium carbonate, 15 pounds of ammonium resinate (dry weight) is then added in the form of an aqueous solution containing 500 pounds of water. 15 pounds of alum

are then added, which immediately reacts with the carbonate to form theoretically 3½ pounds of precipitated alumina. Instead of adding this alum to the beater, the alum solution may first be neutralized with ammonia or other alkali, and the precipitated alumina added to the beater with the size. The hydrated aluminum oxide will combine in the beater with the ammonium resinate to form a compound which coats the first in the beater and which will size the paper when the pulp is dried.

Another method of operation is as fol-

lows:

The carbonate filler, or other filling material, is mixed with water in a tank to a concentration of about 20% solids to which mixture is added an aqueous solution containing ammonium resunate to the extent of about 1 pound of the dry resinate to 100 pounds of filler. To this may be added 1 pound of alum to each 100 pounds of filler along with sufficient ammonia or other alkah to neutralize it

and precipitate the alumina.

This separately treated filling material containing sizing ingredients may be added to the paper stock in the beater, in the Jordan chest, in the machine chest, or at the wet end of the paper machine. This treatment produces a paper containing individually sized filler particles, that is, each particle thereof is coated individually with size. The paper stock in the beater may be sized by the use of ammonium resinate and alumina. If this is done, the result is a paper with fibers and filler particles individually sized with the same sizing materials. Or the paper stock may be first sized with any sodium resinate and sufficient alum to acidify the fibers, whereupon and later, the ammonia sized filler material is added thereto in the beater, machine chest, Jordan, and so forth, whereby a paper is produced having its fibers individually sized by the use of sodium resinate while its filler particles are individually sized with ammonium resinate and alumina. Since the ammonium resinate is somewhat more expensive than sodium resinate, this latter procedure offers some saving in cost over treating both fibers and filler with ammonium resinate.

In general, in the final mixture of paper fibers and filling material, there must be no alkalinity derived from soda. There will be none in the mixture resulting from the practice of this invention because any alkalinity produced by the ammonium resinate disappears on drying of the paper. This produces a neutral

and sized paper.

With the present processes using sodium resinate, it is not possible to fully size a heavily loaded paper containing from 20% to 30% filler even if the filler is not alkaline. By the use of this process, however, any kind of filling ma-terial can be sixed. In order that ammonium resinate may properly function as a sizing material there should always be present enough excess ammonia or other alkali, to form sufficient alumina when reacting with alum to form a resinate of alumina, but it is immaterial how this ammonium hydrate is produced.

Waterproofing Composition II. S. Patent 2,022,405

U. D. 1 access 2,,	
Refined Paraffin Wax	4 lb.
Paracoumarone Resin	2 lb.
White Beeswax	1 lb.
Aluminum Palmitate	4 lb.

The above ingredients being blended together and dissolved in a composite solvent of xylol and carbon tetrachloride in the proportions of about three parts by volume of xylol to one part by volume of carbon tetrachloride, and the amount of solvent being such that about 2% ounces of the above composition is contained in each gallon of solution.

Fireproofing Materials French Patent 774,089

An antiseptic fireproofing composition for wood, paper, etc., contains, e.g., ammonium orthophosphate 5 grams, sodium tetraborate 2.5 grams, and ammonium chloride 2.5 grams.

Exterior Primer 67 lb. 33 lb.

Pigment Vehicle

Pigment: Titanox B White Lead Asbestine Litharge	(Carbonate)	37.1. lb. 37.1 lb. 24.8 lb. 1.0 lb.
Vehicle:		

Archer-Daniels No. 635 lb. Mineral Spirits lby *VM-1367 2% Liquid Cobalt Drier 2, lb.

lb.

boiled oil.

VM 1367: Heat 15 gal. china-wood oll with 75 fb. low acid ester gun to 565° F Remove frozaga and let rase to 585° F, hold for 5 minutel and check with 25 lb. ester gun. Thin at 400° F, with 15 gal. mineral spirits.

Painting Primer German Patent 608,738

Zinc Oxide		30	
Ochre		30	
Linsed Stand Oil		* 14	
Linseed Oil Varnish		21	g.
The above is thinned	with:		
Linseed Oil Varnish	. 1/2		g.
Benzine		9	g.

Exterior Wood Primer

Pigment	()() 11/1
Velucio	34 lb.
Pigment:	
Titanium-Barium Pigment	34 lb.
White Lead (Carbonate)	26 lb.
Metronito	40 lb.
Vehicle:	
Bodied Linseed Oil	13 lb.
Blown Lanseed Oil	5 lb.
Raw Linseed Oil	27 lb.
20 gal Ester Gumwood	20 lb.
Oil Varnish	
Mmeral Spirits	32 lb.
Drier	3 lb.

Priming Paint from Hardened Paint German Patent 607,554

Dissolve old paint in f	ollowing:
Butyl Alcohol	50 lb.
Xylol	10 lb.
Benzol	10 lb. 10 lb.
Toluol	5 lb.
Ethyl Acetate Ether	5 lb.

Galvanized Roof Primer

Bry Red Lead Boiled Linseed Oil Turpentine	10 lb. 9½ ga 1% ga ½ ga	l. l.
Drier	78 B	u

Galvanized Roof Finish

Dry Red Lead	5	lb.
*Oarbon Black Paste	31	lb.
Bolled Linseed Oil	01/4	
Turpentine	17/8	gai.
Drier		
*The carbon black paste formula is 180% carbon	black and	in (iii

Paste Paint L, White Lead Basic Carbonate White 28.4 lb. Lead 3.88 lb. Raw Linseed Oil

JU THE CHEMICA	o Folimonatei
Paste Paint TLZ, Titanox-Lead-Zinc Titanox B 9.8 lb.	White Exterior Bakelite Enamel (Yacht White)
Basic Carbonate White Lead 7.6 lb.	Pigment 40 lb. Vehicle 60 lb.
Zinc Qxide, Lead-Free 4.33 lb. Raw Linseed Oil 3.88 lb.	Pigment:
Spot Priming Paint	Basic Carbonate White Lead 40 lb. Titanium-Barium Pigment 40 lb.
Paste Paint TLZ (above) 1 gal.	Titanium Oxide 20 lb.
Raw Linseed Oil 1 gal. Turpentine 0.28 gal.	Vehicle: *Varnish XV-4430 60 lb.
Drier 0.05 gal.	tVarnish XV-5922 20 lb. Mineral Spirits 20 lb.
Under Coat Paint	Drier:
Paste Paint L or TLZ (above) 1 gal.	Lead 2.5 g. per gallon enamel, as
Raw Linseed Oil 0.51 gal.	naphthenato
Turpentine 0.62 gal. Drier 0.04 gal.	Cobalt 0.15 g. Manganese 0.05 g.
200 - 800	*Varnish XV-4430:
Finish Cont Paint	Bakelite Resin XR-2963 100 lb. China Wood Oil 20 gal. Body Q Linseed Oil 30 gal. Lead Accitate 2 lb
Paste Paint L or TLZ (above) 1 gal.	Body Q Linseed Oil 30 gal. Lead Acetate 2 lb
Raw Linseed Oil 1 gal.	Mineral Spirits 34 gal Dipentene 55 gal.
Turpentine 0.12 gal.	Procedure:
Paint Drier 0.06 gal.	Place the Bakelite, the China wood oil and
Tropical Roofing Paint	Place the Bakelite, the China wood oil and 10, gallous of the lineed oil in the kettle. Heat to 500° F. in one hour. Add the remaining 20 gallous of lineed oil. The temperature will drop to about 450° F. Reheat to 520° F. Add to lead acctate. Cool gurkly should be supported to 450° F, and the wide of the water syring to 450° F, and the wide of the supported to
Paste White Lead 100 lb.	520° F Add the lead acctate. Cool quickly with the aid of water spray to 450° F, and
Non-setting Red Lead 10 lb. Lamp Black in Oil 1/2 lb.	thin with the mineral spirits. †Varnish XV-5922:
Raw Linseed Oil 3 gal.	Bakelita Resin XR-2963 100 lb
Boiled Linseed Oil 1 gal. Turpentine or White Spirit 1/2 gal.	China Wood Oil 7 5 gal. Body Q Linseed Oil 2 5 gal. Lead Acctate 2 5 lb.
Terabine Driers 1 pt.	Lead Acctate 2.5 lb. Lead Carbonate 1.25 lb.
A proportion of hard drying outside	Mineral Spirits 15 gal. Procedure:
quality varnish may be added if desired. Thin out this paint to the desired con-	In 50 minutes heat the Bakelite and China
sistency with equal parts of raw linseed	minutes raise the temperature to 540° F.
sistency with equal parts of raw linseed oil and turpentine. Where the paint must be cheapened, barytes, china clay,	In 50 minutes heat the Bakelite and China wood oil to 450° F. In an additional 18 minutes raise the temperature to 540° F. Add the linseed oil and the driers. Let the temperature drop to 450° in about 20 minutes and the second of the comperature drop to 450° in about 20 minutes.
slate powder, or othre is incorporated as	utes, and thin with the mineral spirits.
an extender.	
Priming Structural Paint	Lead Titanate Exterior Paints
Formula No. 1	Formula No. 1
Dry Basic Lead Chromate 15½ lb.	Lead Titanate 1000 lb. Raw Linseed Oil 252 lb.
Raw Linseed Oil 5 pt. Turpentine 2 gills	China Wood Stand Oil 28 lb.
Liquid Drier 2 gills	Lead-Manganese-Cobalt Drier 8 lb. Mineral Spirits 42 lb.
No. 2 Dry Basic Lead Chromate 151/2 lb.	No. 2
Boiled Linseed Oil 5 pt.	Lead Titanate 400 lb.
Turpentine 1 pt.	Basic Carbonate White
These paints weigh about 21 pounds per gallon and the non-volatile portion	Lead 400 lb. Asbestine 100 lb.
contains about 30% by volume of pig-	Bilica 100 ib.
ment.	Raw Linseed Oil 382 lb.

China Wood Stand Oil	52	lb.
Cobalt Naphthenate	10.8	lb.
Mineral Spirits	65.9	lb.
No. 3		
	400	lb.
Lead Titanate	400	ID.
Basic Carbonate White	400	11.
Lead	400	lb.
Zine Oxide	200	lb.
Raw Linseed Oil	324	lb.
Kettle Bodied Linsced Oil		
(Viscosity Z)	21.6	Ib.
Lead-Manganese-Cobalt		
Drier	20.1	
Mineral Spirits	40.7	lb.
No. 4		
Lead Titanate	400	lb.
Titanox-B	400	lb.
Zinc Oxide	200	lb.
	400	lb.
Raw Linseed Oil	400	10.
Kettle Bodied Linseed Oil	26.4	11.
(Viscosity Z)	-11, 1	11).
Lead-Manganese-Cobalt	25 1	11.
Drier		
Mineral Spirits	50.1	10.
No. 5		
Lead Titanate	200	lb.
Titanox-B	200	lb.
Basic Carbonate White		
Lead	200	lh,
Zinc Oxide	200	lb.
Asbestine	100	lb.
Silica	100	lb.
Raw Linseed Oil	466	lb.
Kettle Bodied Linseed Oil		
(Viscosity Z)	29,6	lb.
Lead-Manganese Cobalt		
Drier	29.2	
Mineral Spirits	58.1	lb.
*This type is of special intere	st for	n or no
a base for house paint tints		

Fire Retarding Interior Whitewash

 Mix about 120 lb. of spent carbide residue with water to a creamy consistency.

Mix 2½ lb. of rye flour thoroughly with ½ gal. of cold water, and then thin with 2 gal. of boiling water.
 Dissolve 2½ lb. of common salt in

3. Dissolve 2½ lb. of common salt in 2½ gal. of hot water.
Mix (2) and (3), then add (1), and

stir until well mixed.

Exterior Weatherproof Whitewash Formula No. 1

- Mix about 120 lb. of spent carbide residue with water to a creamy consistency.
- Dissolve 2 lb. of common salt and 1 lb. of zinc sulphate in 2 gal. of boiling water.

Provide 2 gal. of skimmed milk.
 Pour (2) into (1), then add (3), and stir well.

No. 2

- Mix about 15 lb. of spent carbide residue to a creamy consistency with water.
- 2. Dissolve 1 lb. of carbonate of soda in ¼ gal. of boiling water.
- Soak in cold water for at least 8 lir. ¼ lb. of common glue and 1 lb. of rice flour; and then thoroughly dissolve the glue mixture in ¼ galmere water in a double boiler. Mix (1) with (2), then add (3).

No. 3

- Mix about 12 lb. of carbide residue to a creamy consistency with water.
- 2. Dissolve 4 oz. of white rosin in 12 fluid oz. of boiled linseed oil.
- Beat 6 lb, of whiting in 1 gal, of skimmed milk.
 Mix (2) with (1) while hot, add (3).

Hints for Special Uses

Alum added to whitewash prevents its rubbing off. Flour paste will also prevent rubbing off, but when this is used, zine sulphate must be added as a preservative.

Molasses causes lime to penetrate wood and plaster better. One pint of molasses to 5 gallons of whitewash is generally considered sufficient. A solution of siliente of soda or water glass, one part to ten purts of whitewash, makes what is commonly referred to as a "fire-proof cement" of whitewash.

By adding I pound of cheap bar soap dissolved in I gallon of bothing water, to every 5 gallons of whitewash, a more or less gloss finish can be obtained.

- A fire retardant whitewash, of a type nsed extensively by the U. S. Lighthouse Board, is made according to this formula:
 - Mix about 60 lb. of spent carbide residue with water to a creamy consistency.
 - 2. Dissolve 1 peck of salt in warm water.
 - 3. Add (2) to (1) and mix.
 - 4. Boil 3 lb. of ground rice in water to to a thin paste.
 - 5. Dissolve 1 lb. clear glue in hot
 - 6. Provide 1/2 lb. of powdered Spanish whiting.
 - 7. Mix (4), (5), and (6) together and add to mixture (3). Mix well and let stand for several days.

Keep the wash thus prepared in a kettle or portable furnace, and when used put it on as hot as possible with a painter's brush or whitewash brush.

Cold Glaze for Wall	Tiles
Lacquer Base	
a. Shellac	8 oz.
Turpentine, Thick	5 oz.
Alcohol	35 oz.
b. Sandarac	14 oz.
Turpentine, Thick	6 oz.
Alcohol	35 oz.
Mix 10 oz. of a with	
12 oz. of b	

To this lacquer base add finely powdered pigments, as to color

(Black)

Ultramarine or Paris Blue	(Blue)
Chrome Yellow Zinc Yellow or Ochre	(Yellow)
Chrome Green	(Green)
Chrome Red or Cinnabar	(Red)

Lamp Black

(White) Lithopone (Grind Pigment with a small part of the lacquer solution; thin later with the rest to needed consistency.)

Floor Finish

(Permanent, Scratch-free) Clear (Natural) Finish: Formula No. 1

Castor Oil 1 qt. Boiled Linseed Oil ½ gal. Paraffin Wax 3¼ lb.
Boiled Linseed Oil ½ gal.
Paraffin Wax 3¼ lb.
High-Flash Naphtha 3 qt.
Gasoline 1½ gal.
Varnolene 1 gal.

Mix the oils and wax and heat until the wax is molten. Add the varnolene, naphtha and gasoline slowly in the order mentioned.

No. 2

Dark Finish

Castor Oil	1	qt.
*Gilsonite Cook	1	gal.
Paraffin Wax	3	lb.
High-Flash Naphtha	1	qt.
Gasoline	11/2	gal.
Varnolene	1	gal.

Heat oil and wax until molten, add the gilsonite cook and proceed as above.

*Gilsonite Cook:

Gilsonite		5	lb.	
Kellogg Varnish Oil		11/2	ga	ı.
High Flash Naphtha		1 1/4	ga	J.
Heat gilsonite and oil to 270° (1	(520)°	F.)
Let cool and thin with naphtha.	Ψ.			

Any shade may be obtained by inte-mixing clear and dark finish. Apply b flowing on the freshly scraped floor distribute and rub in lightly with rag-Permit to dry for at least 48 hours. Thi finish actually impregnates the floor an will not wear off. It has a velvet shee and a slight slip, is easy to keep clea. and is very resistant to moisture.

Varnish for Naval Aircraft

raimen tot mavat	Allera	
faterials:		
Bakelite BR-254	50	lb.
Bakelite XR-4036	50	lb.
Castor Oil (Refined)	4.33	lb.
China Wood Oil	33	gal.
Mineral Spirits	27	gal.
Xylol	4	gal.
Dipentene	4	gal.
Lead Cobalt Manganese		_
Naphthenate Driera		

Procedure:

Heat the oil and the Bakelite resins together to 310° F. in 25 minutes, and hold at that temperature for half an hour. Heat to 450° F. in 20 minutes and hold for 20 minutes. Remove from the fire, add the thinners, the castor oil and sufficient drier to give 12 grams cobalt, 15 grams manganese and 160 grams lead as metal.

Airplane Varnish

The naval aircraft factory has developed a formula for satisfactory bituminous varnish which is used for airplane hulls or other parts exposed to salt water or salt spray. This formula is as follows:

Aluminum Powder	2 lb.
Bituminous Primer	1 gal.

Coating for Aluminum or Brass Nitrocellulose 5 g. 55 cc. Amyl Acetate Alcohol 40 cc.

Aluminum Powder Paste U. S. Patent 2,002,891

Aluminum, Flaked	58	oz.
Stearic Acid, Powdered	1	oz.
Aluminum Stearate	1	oz.
Naphtha	40	0Z.
Grind together until homoge	neou	18.

Preparing Aluminum for Enamel

The best method of cleaning aluminum castings, so the finish will adhere tenaciously, is to use the sandblast. Smooth

aluminum surfaces are of such character that an ordinary first coat of finishing material will not adhere to them satisfactorily, even when they are clean. The sundblast will leave the surface slightly etched and will aid the first coat in sticking to the metal permanently.

If sandblasting is impractical, about all that can be done is to thoroughly wash the castings with naphtha or some other solvent for grease, and dry them thoroughly with clean cloths.

In other instances it may be satisfactory to bake the castings for a short time at 400 or 500° F., just before finishing them, to burn off any oil or grease. It is not advisable to use caustic cleaning solutions with aluminum, because the metal is so easily attacked and dissolved by this chemical.

Another method is as follows: Immerse them in a 20% solution of acetic acid until all oil and grease is removed or neutralized. Then rinse in a vat of clear hot water and allow castings to drain and dry. Do not wipe them. Spray or brush as soon as the moisture has disappeared.

Bronzing Liquid

Celluloid Scrap	3	07.
Amyl Acetate	12	oz.
Benzine	28	οZ.
Denatured Alcohol	24	07.

This solution is mixed with sufficient dry gold bronze to make a smooth working paint and the resulting paint must be used at once as it is apt to turn greenish and thicken to a jelly on standing.

Bronze Painting Tinctures

		(Water		90	oz.
А.	a.	{ Water } Alcohol		10	oz.
			 10:		

b. Isinglass or Mirror as desired Gelatin

Add to this colloidal solution with stirring:

c. Bronze Powder sufficient to suit. B. for a and b take:

10 07 Potash-Water Glass Gum Arabic 10 oz. 40 oz. Water

C. or Thick Gum Arabic Solution with a little ox gall.

Paints for Copper

Copper, brouze, or brass gutters and flashings, as well as copper or bronze screening, are apt to cause bad yellowishgreen stains on light- or white painted

houses, owing to the washing off of corrosion products. Exposure tests indicate that one of the best ways to paint copper or bronze surfaces is to wash off any grease, using gasoline or turpentine. The surface should be roughened slightly with sandpaper, and a priming coat composed of 112 to 2 pounds of aluminum powder to 1 gallon of aluminum mixing varnish applied, followed by the desired color cont. Weathered copper or bronze screening should be thoroughly dusted, and then given two conts of a thin black paint. Some of the best grades of black auto top diessings, which are free from asphalt, but are essentially thin, water resistant, carbon black enamels, make excellent sercen enamel.

Cable Lacquer British Patent 397.554

12	07.
12	oz.
50.2	07.
10	02.
10	07
5	02.
	0.8 50.2 10

Electrolytic Condenser Conting British Putent 419,927

137.8 cc. Acetone Amyl Acctate 125 0 ec.

Phenol Formaldeliyde Resin 399 g. Graphite (99%) This is baked on aluminum for 21 hours at 100° C. and 2 hours at 170° C.

Electrical Wire Lacquer British Patent 410,576

Cellulose Acctate Tetrachlorethane 100 oz. Alcohol 20 oz. Tracetin

Adhesiveness may be increased by incorporating tale and opacity by zinc oxide.

Wash for Galvanized Iron before

Painting a Denatured Alcohol 60 ft nz. 30 fl. oz. Toluol' 5 fl. oz. Carbon Tetrachloride Commercial Concentrated 5 fl. oz. Hydrochloric Acid

b. Copper Arctate 6 02. 1 gal. Water

c. Copper Nitrate Crystals 2 oz.
Copper Chloride Crystals 2 oz.
Ammonium Chloride
Crystals 2 oz.
Commercial Concentrated

Solution a will cut grease as well as etch. If the metal is not free from grease, solutions b and c must be preceded by a grease-removing operation.

Treatment of Galvanized Sheets for Painting

A simple and inexpensive way to treat new galvanized sheets before painting is to use ordinary vinegar, either sponged or brushed on. Vinegar rather thoroughly removes the sliek film usually found on newly galvanized sheets. It does not, however, etch the surface like some other treatments. After the vinegar has been applied and allowed to remain on the sheets for five minutes or so, it should be wiped and then the surface of the sheet allowed to completely dry before paint is applied.

Another somewhat similar treatment is the use of two or three per cent acetic acid solution at a temperature of about 130° F. If it is possible to dip the sheets, or articles made from the sheets, in this solution, allow them to remain there for about ten or fifteen minutes. After removal, they should be thoroughly rinsed and allowed to thoroughly dry.

Still another, even more practical, although perhaps a little more costly, method of obtaining a clean and etched surface is to apply, with an oil-free brush, and allow to remain for about ten minutes, an acidified solution made up as follows:

Denatured Alcohol 50 fl. oz.
Toluol 35 fl. oz.
Hydrochloric Acid 5 fl. oz.

This solution should be prepared only as required for immediate use. After the reaction is complete and the surface is thoroughly dried, wash or rinse with clean water to remove any soluble salts that may have formed. Then, allow the sheets to thoroughly dry again before applying paint. This treatment is especially effective if the procedure outlined above is carefully followed.

It should be particularly noted that with each of the three methods outlined, it is important that the galvanized surface should be thoroughly dry before painting. A film of moisture between the paint and sheet would cause very poor adherence.

Painting Galvanized Iron

Excellent paint adherence on galvanized surfaces may be obtained by cleaning with the following solution:

65	lb.
35	lb.
5	lb.
10	lb.
	35 5

This treatment should be followed by a cold rinse after the material has dried.

Lacquer for Hot Water	Containers
Lacquer Linseed Oil	250 g.
Milori Blue	15 g.
Gilsonite	120 g.
Albertol Resin (116° mp	o.) 40 g.
Thick Linseed Oil	40 g.
Manganese Hydroxide	2.5 g.
Cobalt Drier	1.25 g.
Toluol	500 g.

Iron "Lacquer"

zion zaucquei	
Gilsonite Asphalt	20 kg.
Manila Copal	5 kg.
Lampblack	3 kg.
Toluol	50 kg.

Iron Protective Paint

Formula No. 1

Lampblack (Ground in Oil)	90.7	oz.
*Asphalt Varnish	68.1	oz.
Linseed Oil, Raw	68.1	oz.
Japan Drier	2.0	oz.

No. 2

110. 2		
Lampblack	27	oz.
Silica	58	oz.
Red Lead	10	oz.
Graphite	5	oz.
*Asphalt Varnish	Suffic	cient
Grind together until smoot	th.	
*Turpentine Asphalt in Linsced Oil	1	part part

Primers for Light Metal Alloys

Owing to high coefficient of expansion and contraction with temperature changes, a primer is needed that will be sufficiently flexible not to be ruptured by expansion and contraction. A zinc chromate paint is recommended for this purpose, a specimen formula being:

ose, a specimen ronnula being	۶٠	
Zinc Chromate	40	lb.
Neutral Red Oxide of Lead	80	lb.
Boiled Linseed Oil	60	lb.
Pure Turpentine	16	lb.
Strong Japan Driers	4	lb.

Another priming paint found to be satisfactory is made from:

Dry Lampblack 65 lb. Linseed Oil 15 lb. Pure Turpentine 10 lb.

Driers according to type and quality.

The primer should be allowed 50 to 60

The primer should be allowed 50 to 60 hours to dry and harden before applying subsequent coatings.

Polished Metal Lacquer

Nitrocellulose Wet (15 20 sec.)	10	g.
Rezyl No. 468-2 (50% So-		
lution)	10	
Dibutyl Phthalate	2	
Butyl Acetate	10	
Butyl Alcohol	- 8	g.
Butyl "Cellosolve"	10	g.
Toluol	35	g.
Xylol	15	g.

Preparing Magnesium Alloys for Painting

To prepare the surface of magnesium alloys so that paint will adhere, it is recommended that the alloy be first immersed in the following:

Sodium Diehromate 1.5 lb. Concentrated Nitric Acid 1.5 pt. Water 1.5 up. 1.5 pt. 1 gal.

In a new solution, only 15 seconds are needed. This time increases to two minutes for an old solution.

After rinsing and drying, the proper primer should be used, containing inert pigments or, for example, zinc chromate. For interior work, a minimum of two coats (total) paint should be used; for exterior work, a minimum of four coats.

Care and Preservation of Bronze Statues

Statues, tablets, medals, especially those standing in the open, require careful treatment and protection from the conditions tending to their corrosion. Of cleaning reagents, water only is permissible with, perhaps, a small quantity of soap extract. Bronze which has become black by long exposure may be restored to its original gold color by washing with water to which a little ammonia is added, using a brush with bristles, no wire brush.

As protective coating, a mixture of beeswax and turpentine is considerable protection to bronze from atmospheric attack and gives a pleasing appearance, besides drying rapidly. Applied three times a year it will safeguard a statue to a

high degree from corrosion and deterioration even in an exposed position. A mixture of hinolin and paraffin is not quite as good as it does not dry as rapidly and is therefore hible to collect dust.

Heatproof Rust Protective Contings

Kerosene and pitch cannot be used as buiders as they become too soft even at 150 200° C. Natural asphalts, although brittle, give protection up to 250° C, acetyl cellulose up to 100° C. Only lean, not fatty brading agents rich in resms, should be used for such paints. As at 400° C, almost all binding agents are entirely dismitgrated, the residues of the agents must be such that they leave a continuous, well adhering coat on the metal to be protected. Durophen, aluminum bronze, zinc dust with binders of this type give good results. Heatproof paints should never be sprayed on, as they have the tendency to spall off later, but brushed on, except zinc dust which may also be sprayed.

Rust Proofing

A good protective coat for metal articles during storage and transit is unde by brushing on a solution of landing metals white spirit or solvent naphtha. Equal weights solvent and landin seem satisfactory and there is not much to choose between the two solvents. If a rather harder film is wanted, up to 5% ceresin wax can be added in the case of naphtha solutions; in the case of white spirit up to 10 per cent parafflin wax or up to 3 per cent ceresin wax. It is recommended that the white spirit be of the BESA, standard, i.e., B.P. 160° to 110° approximately and is to the landing the results of practical tests show little difference between widely different grades.

7.8 lb. lanolin in 1 gal. white spirit give 1.9 gal. solution. 8.3 lb. lanolin in 1 gal, solvent naphtha give 1.9 gal. solution.

Crystal Coating on Steel

Orjana Couring	5 OIL DUCCE	
Sodium Nitrate	3	lb.
Manganese Dioxide	3	lb.
5% Sulphuric Acid 8	Solution 100	gal.

Protective Coating for Structural Steel

Coal Tar Pitch	62.5 lb.
Benzol	25 lb.
Aluminum Bronze	12.5 lb.

Priming Structural Paint (Red Lead)
Formula No. 1	inca incaa,
Dry Red Lead	20 lb.
Raw Linseed Oil	5 pt.
Turpentine	2 gills
Liquid Drier	2 gills
No. 2	
Red Lead Paste in Oil	20 lb.
Raw Linseed Oil	3 pt.
Turpentine	2 gills
Liquid Drier	2 gills
Finish for Steel Surfa	aces
Tornesit	20 g.
Linseed Oil, Crude, Boiled	10 g.
Indian Red	20 g.
Xylene	30 cc.
High-Flash Naphtha	40 cc.
First Coat Structural Steel	Protective
Blue Lead, Paste in Oil 1	00 lb.
Raw Linseed Oil	2¾ gal.
Turpentine or Mineral	
Spirits	1% gal.
Drier	1/4 gal.
Top Coal Structural Steel	l Paint
Pigment:	
C.P. Chrome Orange	90 lb.
Magnesium Silicate	10 lb.
Vehicle:	
Raw Linseed Oil	80 lb.
Spar Varnish	10 lb.
Liquid Paint Drier	10 lb.
Paint:	70.11
Above Pigment Above Vehicle	70 lb. 30 lb.
Above venicle	30 10.
Red Lead for Brid	• ·
Red Lead	40 lb.
Iron Oxide (95%)	40 lb.
Stand Oil	90 lb.

Tin Can Coating U. S. Patent 2,009,776

12 lb. 20–40 lb.

1 lb.

Raw Linseed Oil

Turpentine Cobalt-Manganese Drier

A coating dough for producing a coating material comprises & mixture of 100 parts by weight of rubber solution containing approximately 30 parts by weight of rubber, approximately 15 parts by weight of adhesive ester gum, approximately 3 parts by weight of liquid petrolatum, and approximately 100 parts by weight of zinc oxide.

Tin Lithographing Varnish

Typical construction of this class of product is represented by the following formulae: 54 gal. pale amberol varnish, 34 gal. gum solution, 22 gal. pale mixing varnish, 8 lb. of white vaseline warmed and reduced with 2 gal. of mineral spirits.

The first component of the above blend, is -135 lb. amberol F7 light, 15 lb. WWX Rosin, 34 gal. pale China wood oil, 1½ gal. "Superior" linseed oil, 6 gal. bodied linseed (1½ hrs. at 600° F.), 10 lb. fused lend resinate, 1 ounce cobalt acetate, 8 gal. gum turpentine, 65 gal. mineral spirits.

The second component is a solution of ester gum in mineral spirits, using 12½ lb. of gum to each gallon of solvent.

The third component is 50 lb. ester gum, 3 lb. fused lead resinate, 10 lb. WWX Rosin, 50 gal. pale China wood oil, 50 gal. mineral spirits.

"Tornesit" Paints

First, a base solution is prepared, consisting of 33½ per cent Tornesit and 6635 per cent high-flash naphtha. To effect solution is a matter of a very few minutes, if the "Tornesit" is added to the solvent.

Second, a concentrated gum solution is made when required.

Third, the pigments are ground in the plasticizer, or if it is insufficient, some of the "Tornesit" base solution is used. Fourth, if a brushing paint is required,

Fourth, if a brushing paint is required, the base solution is thinned to a ''Tornesit'' content of 21 per cent to 22 per cent by the addition of a solvent mixture consisting of two parts high-flash naphtha and one part xylol. If a spraying composition is desired, the base solution is thinned with toluol to a ''Tornesit'' content of 11 per cent to 12 per cent. It is advisable to ship even spray paints with a brushing viscosity and send the thinner separately. This helps to keep the pigments in good suspension.

Finally, the gum solution and pigment paste are added to the reduced solution and the mixture is stirred.

"Tornesit" paints may be applied by spraying, dipping, flowing, or brushing. A good film can be obtained by any of these methods.

Following is a brief outline of procedure to be followed, to obtain most satisfactory results in spraying and brushing:

Spraying

"Tornesit" solutions can be sprayed, producing a hard, durable, evenly distrib-

uted film. With present equipment, the spraying viscosity is 40 centipoises, which is somewhat lower than the 75 centipoise spraying viscosity of lacquers.

If the "Tornesit" concentration is

If the "Tornesit" concentration is kept below 12%, no difficulty will be countered from "spider-webling." By the addition of softening agents, gums, and pigments, the solids content will be increased 30-40 per cent, depending, of course, on choice of ingredients.

Brushing

Brushing paints with as high as 57 per cent solids have been applied successfully. For this purpose, a working viscosity of about 250 centipoises is recommended. In brushing "Tornesit" paint, the surface should be well covered with a full brush, avoiding going over the painted area any more than necessary because of the rapid drying of the product. When bothed tung oil is used as the plasticizer in the priming coat, a second coat may be applied to an interior surface after six to eight hours. On exterior work, three to four hours is an ample drying period with the same priming coat.

"Tornesit" Paints

A formula used successfully on tank cars, structural steel and smular surfaces not subject to immersion contains Tornesit plasticized with a drying oil. China wood oil must be properly boiled to avoid wrinkling when a second coat is applied, but no wrinkling occurs with Inseed oil. When properly formulated, "Torneat" paint has good adhesion to metal. Examples of primers having good adhesion are:

Formula No. 1 No. 2

101	munt Ivo	1 1.0
"Tornesit"	20 oz.	20 oz.
Heavy-Bodied Raw		
Linseed Oil	10 oz.	10 oz.
Cumar PlO		5 oz.
Iron Oxide	20 oz.	20 oz.
Silica	30 oz.	
Xylol	70 oz.	70 oz.
A finish coat use	ed succes	sfully on
"Tornesit"		20 oz.
Heavy-Bodied Raw		
Linseed Oil		10 oz.
Indian Red		20 oz.
Xvlol		30 oz.

Formulæ containing improperly-bodied oils do not have good alkali resistance, but to withstand immersion in aqueous media, particularly those containing alkalies, formulæ such as the following have been quite successful:

High-Flash Naphtha

	Formula	No. 1	No. 2
"Tornesit"	20	OZ.	20 oz.
Methyl Abictate	12	oz.	16 oz.
Cumar V			24 oz.
Indian Red	25	()Z.,	-
Titunium Dioxie	le —		40 oz.

Finishes made to the foregoing formula containing iron exide have withstood immersion in 5 per cent caustic soda for two months and in 5 per cent hydrochloric acid for three weeks, the use of iron in the pigment probably reducing resistance to hydrochloric acid.

Pliolite Varnish (Paper	Coating)
Pliolite Resin	15 g.
Ester Gum Solution (4 # cut)	10 g.
Tricresyl Phosphate	5 g.
Toluol	70 g.

Paper Enamel

U. S. Patent 2,000,453

Glue	20	oz.
Anmonum Hydroxide	2	02.
Alcohol	4	07.,
Chromic Acid	11/2	02.
Water to make	1	gal.

Moisture Proof Paper Lacquer British Patent 412,687

Ozokerite	1-2 oz,
Dibutyl Phthalate	25-50 oz.
Nitrocellulose	50-75 02.
Lacquer Solvent	to suit

Paper Watermarking Fluid

O. o. Patent synal, ter		
Canada Balsam	8-20	lb.
Turpentine	5-17	lb.
Colorless Mineral Filler	8-25	lb.
Castor Oil	12 - 30	
Borax Solution (1%)	sufficient	to
amplety above liquid	R.	

Water to thin to working consistency.

Rubber Paints

British Patents 407,038 and 417,912

Preparation of Solution "B"

Raw crepe rabler is masticated on a rubber mill, using warm rollers, until the rubber runs coherently round the rollers. Keeping the rubber still milling, 2½ per cent of cohalt linoleate (6 per cent metallic cobalt content) is then added, when the cobalt linoleate is completely dispersed in the rubber, the mixture is

taken off the mill and immediately transferred to a solution mixer, and churned up with an equal weight of white spirit, until a homogeneous mass is formed. This is then poured into drums and is ready for use. The solution should not be kept at a lower concentration than 50 per cent, as there is a tendency for thinner solutions to reduce still further in viscosity and to lose some of their properties.

Preparation of Paint

To prepare a paint, the rubber solution is mixed to the oil with sufficient white spirit to make a medium, which when mixed with the necessary pigments, will form a suitable paste for grinding. Any of the usual pigments and fillers can be incorporated. The ground paste is then thinned with further white spirit to brushing consistency.

brushing consistency.

As examples of up-to-date formulæ for rubber paints, the following are suggested:

ested:			
Flat Paints			
Formula No. 1			
Lithopone	150 lb.		
Yellow Ochre	1.5 lb.		
Middle Chrome Yellow	1.5 lb.		
Solution "B" (above)	20 lb.		
Boiled Oil	10 lb.		
White Spirit	30 lb.		
No. 2			
Lithopone	65 lb.		
Titanium White	65 lb.		
Asbestine	15 lb.		
Solution "B"	20 lb.		
Stand Oil	10 lb.		
Liquid Driers (Lead .033;			
Cobalt .004)	1 lb.		
White Spirit	30 lb.		
No. 3			
Ultramarine Blue	75 lb.		
Asbestine	25 lb.		
Solution "B"	28 lb.		
Boiled Oil	14 lb.		
White Spirit	60 lb.		
No. 4			
Lithopone	100 lb.		
Solution "B"	20 lb.		
Ester (lum	10 lb.		
White Spirit *	50 lb.		
No. 5			
	150 lb.		
Lithopone Stand Oil/Wood Oil (3/1)	10 lb.		
Solution "B"	20 lb.		
Liquid Driers	20 lb. 1 lb.		
White Spirit	30 lb.		
······································	00 104		

Ready-Mixed Gloss Paints			
No. 6			
Zinc Oxide	100 lb.		
Pale Boiled Oil	62.5 lb.		
Solution "B"	25 lb.		
Terebene	2 lb.		
White Spirit	10 lb.		
No. 7	10 10.		
Zinc Oxide	50 lb.		
Titanium White	50 lb.		
Pale Boiled Oil	62.5 lb.		
Solution "B"	25 lb.		
Terebene	2 lb.		
White Spirit	10 lb.		
No. 8	10 10.		
Lithopone	80 lb.		
Zine Oxide	20 lb.		
Pale Boiled Oil	30 lb.		
Solution "B"	15 lb.		
Terebene	1 lb.		
White Spirit	20 lb.		
No. 9			
White Lead	100 lb.		
Pale Boiled Oil	30 lb.		
Solution "B"	12 lb.		
Terebene	1 lb.		
White Spirit	6 lb.		
	•		
Cheap Rubber Paint			
Molten Rubber	100 oz.		
White Spirit	100 oz.		
Terebene	12 oz.		
Cobalt Tarchone	19 00		

Molten Rubber	100 oz.
White Spirit	100 oz.
Terebene	12 oz.
Cobalt Terebene	12 oz.
Red Ochre	100 oz.
The defects of molter	rubber as a
aint vehicle may be obvis	ated by using it
n conjunction with oil.	That is to say,

paint vehicle may be obvinted by using it in conjunction with oil. That is to say, the varnish is made up partly of molten rubber and partly of Inseed oil. A paint made up on a varnish of this description prepared by "cooking up" the oil and rubber together (in the proportion of 50/50) in the presence of driers and thinning with solvents—appears to have good ageing properties and to yield a film which does not readily crack.

* Molten Rubber Varnish Terebene	140 oz. 5 oz.
Red Ochre	100 oz.
* Molten Rubber Linseed Oil + Driers White Spirit	35 os. 35 os. 70 os.

Rubber Water Paint

Glue Solution	25 oz.
Casein Solution	25 oz.
Latex	30 oz.
Lithopone	100 oz.
Drying oils con if desired	ha inaan

Drying oils can, if desired, be incorporated with the above, and for some

-			
purposes are an advantage	. but tend to	Cobalt Linoleate	1 oz.
discolor the paint more ra		Zinc Oxide	100 oz.
Distempers can also be		White Spirit	40 oz.
prepared by using a rubbe		. (Rubber Resin	1025 08
used for the oil paints).	The solution	(White Spirit	King or
readily emulsifies with a			
with which the pigments		* * * *	
	vitil be tilled	Rabber Lacquer	
porated.	1 0	Nitrocellulose, Wet	
The following is an ex-	ampie of this	(15/20 Sec.)	5.0 g.
type of distemper:		Staffey Oil (Plusticizer)	
Glue Solution	20 oz.		2.5 g.
* Rubber Solution	16 oz.	Ethyl Acetate	10.0 g.
		Butyl Acctate	10 0 g.
Water	25 oz.	C.D. Alcohol	10.0 g.
Lithopone	100 oz.	Toluol	62.5 g.
(Milled Crape	8 oz.		
* Milled Crepe Cobalt Linoleate White Spirit	0 2 02	Rubber Repairing Lac	coner
White Spirit	8 oz		4
		(For Galoshes)	
20.11 77 17 17		(Alcohol	240 cc.
Rubber Frosting V	arnisa		1 0
The addition of rubbe	r solution to	" Nigrosin (Alcohol Solub	
china wood oil gives a fro		(Nigrosin Base BT	50 g.
		b. Benzol (90%)	180 cc.
which will give the desire			200 ec.
more regular manner that		l Acetone, Technical	- m.
wood oil is used alone. Th	e rubber solu-	To 350 cc, of this dyestuff	solution add
tion containing cobalt line	oleate is suit-		
able for this purpose.		Xylene, Technical	350 cc.
	0.0	Vmapas BP. 50T	300 g.
* Rubber Solution	20 07.	Mrx thoroughly, filter throa	ach a cauza
China Wood Oil	10 oz.		Pur to Burner
Terebeno	1 oz.	filter.	
White Spirit	10 oz.		
		Black Rubber Tire P	aint
* Milled Crepe * Cobalt Linoleste	10 oz. 0 25 oz	Rosin	3 kg.
White Spirit	10 oz	Turpentine	3 kg.
(white spirit	10 02		
		Shellae	12 kg.
Rubber Flat P:	int	Sandarae	6 kg.
Rubber Solution	51 oz.	Alcohol	9 kg.
	01 02.	Turpentine, Venice	3 kg.
Milled Crepe]	Carbon Black	to suit
Rubber 10	07.		
	0Z. (
White Spirit 40		Elastic Covering	
•			249
Stand Oil	10 07.	French Patent 762,:	345
Cobalt Linoleate	0.25 oz.	Viscose	15-30 g.
Lithopone	150 oz.	Rubber Latex	50-80 g.
White Spirit	40 oz.		70
waite opini		Casein	70 g.
		Water	45 g.
Rubber Gloss Pa	int	Sodium Silicate (36° Bé.)	
* Rubber Solution	25 oz.	Hardwood Flour	70 g.
	621/2 oz.	Asbestos Fibers	35 g.
Pale Boiled Oil			40-60 g.
Terebene	2 oz.	Ochre, Unculcined	40-00 B.
Zinc Oxide	100 oz.		
White Spirit	10 oz.	Rubber-Asphalt Lace	mer
	containing co.		
Modified rubber solution	containing co	Crepe Rubber (Shredded)	
balt linoleate (us p	reviously de-	Benzol	90-95 oz.
scribed).		Allow to sonk over night	and stir the
Solution as above, after	blowing with	Allow to soak over night and stir the	
		1 *	
air.		Dissolve	
* Milled Crepe (including * 2 1/2 % Cobalt Linoleate,		Gilsonite	30-40 oz.
* {2 1/2 % Cobalt Linoleate,		in	
(White Spirit 12 1/2)	0.5		60-70 oz.
* Rubber Resin Varnish	25 oz.	Benzol	
Stand Oil	50 oz.	Run the rubber solution in	
Terebene	5 oz.	solution slowly while stirring	ζ.
		,	

Linoleum Preservative
Formula No. 1
Linseed Oil (Free from Mucous Substances)
No. 2

45 g. Caoutchouc, Crude, Soft Resin, Coumarone 15 g. Spindle Oil, Refined 940 g.

Melt up together on water bath.

Linoleum Finish

U. S. Patent 1,99	18,927
Glyceryl Phthalate	12.5 lb.
Tolnol	48.1 lb.
Triethanolamine	0.9 lb.
Apply to uncured plastic	linoleum body
ind keep at about 75° C. f	or 14 days.

Eggshell Enamel	
Pigment	50 lb.
Vehiclo	50 lb.
Pigment:	
French Process Zinc Oxide	80 lb.
Celite No. ON-165	10 lb.
Titanium Dioxide	10 lb.
Vehicle:	
Kettle-Bodied Linseed Oil	60 lb.
Mineral Spirits	35 lb.
Liquid Cobalt Drier	5 lb.

Enameling over Varnish

First wash wood work; sandpaper; mix First wash wood work; sanupaper; mx flat paint or enamel undercoat with a little enamel and brush it out thinly. While wet rub with pumice stone and then smooth coating with a brush. Only a small section may be done at a time. If coating sets too quickly add a little linseed oil.

Aluminum Lacquer		
Beckosol No. 1, Solid	100	σ.
Solvent Naphtha	100	g.
Chlorinated Rubber	20	
Xylene	70	
Cobalt-Siccative (1% Cobalt)	5	
This lacquer is resistant to be	nzol.	Ī

Analytical Weight "Lacquer" Bleached Shellac Alcohol, Pure

Put in corked bottle; shake and allow to stand for a few days. Filter through fine filter paper.

Brushing Lacquer U. S. Patent 1.533.616

01 101	,
Alcohol	10 oz
Ethylene Glycol	10 oz.
Amyl Acetate	5 oz.
Butyl Acetate	10 oz.
Ethyl Acetate	15 oz.
Benzol	15 oz.
Toluol	10 oz.
Xylol	10 oz.
Gasoline	10 oz.
Amyl Alcohol	5 oz.
Butanol	5 oz.

Crystal Lacquer

, ,	
Nitrocellulose, Wet (1/2 sec.)	8 g.
Tunguran "A" (Plasticizer)	9 g.
Furfural	12 g.
Butyl Acetate	8 g.
Ethyl Acctate	30 g.
Toluol	33 g.
	•

Lacquer Thinner

Tolucne	50 cc.
Ethyl Acetate	18 cc.
Alcohol	12 ec.
Amyl Acetate	20 cc.

Cellulose Solution No. 1

Nitrocellulose (Dry Weight)		
(½ sec.)	25	g.
Alcohol	10.7	g.
Butyl Acetate	16.1	
Toluene	32.1	g.
Ethyl Acetate	16.1	ġ.
No. 2		_

Mittocentiose vory	weight)		
(1/2 sec.)	0 ,	35.8	g.
Butyl Acetate		24.8	g.
Toluol		24.2	g.
Ethyl Acetate		15.2	g.

Crystallizing Lacquer Thinner

Olybean Ming Zaroques	
Ethyl Acetato	1.5 g.
"Cellosolve"	0.5 g.
"Cellosolve" Acetate	0.5 g.
Methanol	0.5 g.
Toluene	7 g.

If using phthalic anhydride, make up solution in cyclohexanone, if using naphthalene dissolve in toluene. The resulting solution is stirred into the lacquer. Variations are made by using mixtures of both, naphthalene and phthalic anhydride.

Crystallizing Lacquer

Formula No. 1		
Cellulose Solution No. 1		
(see above)	15	g.
Cellulose Solution No. 2	0.5	ø.

ŧ

COATINGS,	PROTECT
Cyclohexanone Ester Gum in Toluol	6.5 g.
(I:I. Weight)	2 g.
Tricresyl Phosphate	0.5 g.
Amyl Acetate	5 g.
Phthalic Anhydride, or Na	ph-
thalene Flakes	4 g.
No. 2	
Nitrocellulose (1/2 sec.)	4 σ.
Nitrocellulose (100 sec.)	
Butyl Acetate	0 ~ 0
Butyl Acetate Ethyl Acetate	9.5 g. 9.5 g.
Cyclohexanone	8 g.
Butyl Propionate	9.5 g.
Toluene	2 g.
Methanol	3.25 g.
Thinner (see below)	9 g.
Ester Gum in Toluol (1:1) 7.5 g.
Phthalic Anhydride or	
Naphthalene Flakes	8.5 g.
The phthalic anhydride is	to be dis-
solved in the cyclohexane	one (heat
gently), then stir solution into	lacquer.
•	Bulbs
Nitrocellulose	20 g.
Butyl Acetate	0.5 g.
Acctone	50 g.
Alcohol	30 g.
Lithopone, optional or other Pigments.	5–10 g.
or other Figments.	
-	
Spirit (Furniture) Lac	
Shellac, Bleached	25 g.
Sandarac	8 g.
Turpentine	4 ec.
Alcohol, Denatured	100 cc.
Man Drink Income	
Floor Paint Lacque	·r
Formula No. 1	100
a. Rosin Wood Oil, Crude	100 g. 60 g.
Linseed Oil	0
(Linseed On	. •
b. Zine White	4 g.
c. Litharge Manganese Oxide-Hydrate	3 g.
	e 0.5 g.
d. Lacquer Benzoline	100 -
(White Spirit)	160 g.
Heat up a together to 18	0-200° C.,
then add b together with lime the oils). Heat up to 290° C	neuran or
the oils). Heat up to 290 C	falls to
the fire. When temperature 250-260° C., add c.	, , , , , , , , , , , , , , , , , , , ,
When cooled, thin with d.	
,	
No. 2	100 m
a. Kopol No. 600	100 g.
Wood Oil, Crude	70 g.
o. Linscen Oil- Standon	30 g.
Thick	no R.

c. Lacquer Benzoline	160 g.
d. Cobalt Siccative, Liquid	
(1% Metal Content)	6-8 g.
Heat a to 280-290° C., ther	"auench"
with b. When cooled to 180	° C., add c.
hen d.	. ,

Floor Lacquer

Copal Ester Linseed Oil"Standoil"	100 g. 70 g.
Lead Manganese Resinate	4 g.
Cobalt, Siccative	1 g.
Thinner	150 g.
Linoleum or Floor Lacq Nitrocellulose, Wet (1/2 sec.) Dewaxed Damar Gum So-	14 g.
lution (4# Cnt) Paraplex 5 B Solution (80%	12 g.
by Weight) (Plasticizer)	12 g.
Dibutyl Phthalate	2 g.
Toluol	15 g.

Hat Lacquer

Mmeral Spirits
Butyl Alcohol
Butyl Acetate
Butyl "Cellosolve"

Use 1.25 gal, of the damar lacquer shown below to 3.75 gal, of the second thinner although other thinners can be used.

A lacquer may be made from damar gum and introcellulose as follows: 12.5 gal, benzene; 12.5 gal, tolloi; 50 lb, 5sec, introcellulose; 10 gal, ethyl accutate; 8.75 gal, butyl acctate; 21.25 gal, dewaxed damar solution.

The yield is 67 gallons of lacquer. Put the five second introcellulose in a 100 gal barrel or drinn and wet it down with the toluol and a low boiling petroleum lacquer thinner. After mixing them, add the citiyl acetate, half acetate, and dewaxed damar solution. Sirr by hand with a wooden stick, or a power stirrer. The dewaxed damar solution is made quickly by grinding to about 10 mesh: 80 pounds of No. 1 Battavia or Singapore damar goin and adding it to—2.7 gallons of ethyl acetate and—6.43 gallons of ethyl acetate and—6.43 gallons of ethyl acetate and—6.43 gallons of 200 proof alcohol, for cutting shellac. After adding the alcohol, a white waxy precipitate will be formed which will take from one to three days for settling out, depending upon the kind of alcohol used.

The lacquer just described is usually thinned with two parts of a suitable thinner to one part of lacquer before dipping hats into it. The hats are put on racks to dry before shaping on the hot block. A very agreeable non-poison-ous thinner is made by mixing: 53% cleaners' naphtha; 15% butyl acetate; 24% No. 1 Special or other similar solvent; 6% butanol; 2% butyl lactate.

Marble Effect Lacquering German Patent 597,114

Marble effects are gotten by applying the following oil coating over a ground coating of lacquer and then spraying on immediately a very thin lacquer.

Paraffin Oil	40 oz.
Toluol	20 oz.
Alcohol	20 oz.
Ethyl Acetate	20 oz.
Pine Oil	20 oz.
Castor Oil	5 oz.

Non-Inflammable Lacquer

Cellulose Acetate	20 g.
Plasticizer	20 g.
Ethylene Dichloride	120 g.
Ethyl Acetate	30 cc.
Alcohol	20 сс.
Methyl "Cellosolve" "Cellosolve" Acetate	20 cc.
"Cellosolve" Acetate	5 cc.

Pavement Lacquers Formula No. 1

14 g.

Manila Copal	30 g.
Linseed Oil	22 cc.
Cobalt Linoleate Drier	1 g.
Benzoline	33 cc.
No. 2	
(Alcohol	40 cc.
a. Alcohol Manila Copal	40 g.
"'Galipot'' in Alcoholic	
b. "Galipot" in Alcoholic Solution (1.5:1)	20 cc.
[Rosin in Alcoholic So-	
o. Rosin in Alcoholic Solution (2:1)	20 cc.
Mix solutions a, b, c.	

Lacquer Plasticizer

Coconut Fatty Acids	2610	lb.
Sulphuric Acid (66° Bé.) Denatured Alcohol Caustic Soda (14° B Caustic Soda (30° B	é.)	lb. gal.

Manipulation:

Rosin, Pale

1. The coconut fatty acids must be saponified by boiling with excess of is governed in the first place by the pro-

strong caustic soda solution (30° Bé. or stronger) and with addition of considerable water after saponification to prevent solidification of the soap.

2. This soap solution is then decomposed with sulphuric acid, the resulting coconut fatty acids (now being free from neutral oil) are washed with hot water. 3. The fatty acids are heating in a lead lined pressure vessel at 20 to 25 pounds pressure with denatured alcohol and sulphuric acid to esterify to the ethyl ester of the mixed fatty acids. This operation is carried on until the free fatty acid test shows only 6-7 per cent, beyond which point it is uneconomical.

4. The remaining free fatty acids are then neutralized with a 14° Bé. caustic soda solution in a steel tank, allowed to settle over night and the mixed esters pumped off from the soapstock to the

still for distillation.

5. The esters are distilled under 25-26 second vacuum at a temperature of 250-425° F. in a steel mill equipped with oil heat or with means for circulating the esters through a direct heater. The condensing equipment is equipped with a sight glass so that the first runs, which are dark in color, may be separated for addition to the next lot of acids to be esterified. When the distillate becomes pale yellow it is suitable for the finished product receiver. The finished product is bleached water white with Fuller's Earth and decolorizing carbon.

Lacquer Thinners Formula No. 1

Ethyl Acetate Butyl Propionate Toluol	25	oz. oz.
No. 2		
Ethyl Acetate	5	oz.
Butyl Propionate	10	oz.
Fusel Oil		0 2.
Toluol		0Z.
Xylol	10	OZ.
No. 3		
Amyl Acetate	20	oz.
Butyl Alcohol	10	oz.
Methyl Alcohol		oz.
Toluol	60	oz.
No. 4		
Benzine		oz.
Amyl Acetate		0 Z.
Butyl Acetate		oz.
Acetone	20	oz.

Lacquer for Synthetic Plastics

The consistency of a particular lacquer

COATINGS, I	ROTECTI	VE AND DECORATIVE	
a l familiaria T	1	Danatural Alaskal	10 oz.
posed mode of application. I	n general,	Denatured Alcohol	25 oz.
spray lacquers contain 12 to 1	4 per cent	Barium Sulphate	25 os.
nitrocellulose; dipping lacque	es contain	Zinc Oxide	20 04.
8 to 12 per cent nitrocellulose,	and brush	No. 2	
lacquers contain 14 to 17 per o	ent nitro-	Cellulose Acetato	15 oz.
cellulose.		Methyl Acetate	5 oz.
Solvent mixtures will also	vary with	Lacquer Solvent	30 oz.
the mode of application. A t	ypical sol-	Barium Sulphate	25 oz.
vent mixture for cellulose lacq	uers com-	Chrome Yellow	25 oz.
prises:			
Lacquer Solvent		No. 3	
	50 oz.	Nitrocellulose	10 oz.
Ethyl Acetate	20 oz.	Ether	15 oz.
Butyl Acetate		Alcohol	25 oz.
Butyl Alcohol	5 oz.	Barium Sulphate	25 oz.
Benzol	25 oz.	Ochre	25 oz.
The following lacquer compo	sitions are	No. 4	
recommended for highly pol-	ished sur-		15 oz.
faces:		Pyroxylin	35 oz.
Formula No. 1		Lacquer Solvent	25 oz.
	40 oz.	Barium Sulphate	25 oz.
Butyl Acetate	10 oz.	Chrome Orange	
Ethyl Acetate	25 oz.	Greater adhesion can be	secured in
Alcohol		above formulæ by addition of	3 % cuter
Benzol	10 oz.	gum.	
Remainder nitrocellulose, in	cluding 10		
nor cont plasticizer (calculat	ed on the	m. c. ! 1	0.0011.07
nitrocellulose) such as dibutyl	or diamyl	Capsule or Tube Scaling L	acquers
phthalate.		Formula No. 1	
No. 2		Celluloid Scrap	15 oz.
Nitrocellulose	8 oz.	Lacquer Solvent	40 oz.
Shellac	5 oz.	Alcohol, Denatured	25 oz.
Plasticizer	2 oz.	Lampblack	20 oz.
	25 oz.	No. 2	
Alcohol	40 oz.		20 oz.
Butyl Acetate	5 oz.	Cellulose Acetate	5 oz.
Butyl Alcohol	10 oz.	Methyl Acetate	50 oz.
Acetone	5 oz.	Lacquer Solvent	25 oz.
Glycol Monoacetate	0 00.	Zinc Oxide	25 02.
		No. 3	
Lacquer Scalers		Nitrocelluloso	15 oz.
Formula No. 1		Ether	22 oz.
	1 04 lb.	Alcohol	38 oz.
Blown Linseed Oil	9 22 lb.	Ultramarine Blue	25 oz.
Nitrocellulose (14 sec.) Wet	1 gal.		
Tulmer	* B		
Lacquer Thinner:		Transparent Tube Lac	quer
Toluol	61 oz.	Formula No. 1	
Butyl Acetate	26 oz.	Celluloid Scrap	20 oz.
Butyl Alcohol	13 oz.	Lacquer Solvent	50 oz.
No. 2		Alcohol, Denatured	20 oz.
	.47 lb.	Butanol	8 oz.
Nitrocellulose (1/2 sec.)	.93 lb.	Soluble Lacquer Color	2 oz.
Nitrocellulose (40 sec.)	.93 lb.	Boluble Dacquer Sales	
Ester Gum	.93 lb.	No. 2	
Calcium Stearate	1 gal.	1	18 oz.
Tummer	* P	Nitrocelluloso	15 oz.
Lacquer Thinner:		Butyl Acetate	68 oz.
Coal Tar Naphtha	60 oz.	Lacquer Solvent	2 uz.
Butyl Acetate	40 oz.	Soluble Lacquer Color	
Dutyl Acctate			
- I I		Lacquer for Tennis Ra	ckets
Sealing Lacquer			33 g.
Formula No. 1		Manila Copal	66 cc.
Celluloid Scrap	10 oz.	Alcohol (93-95%)	
Lacquer Solvent	30 oz.	Linseed Oil Fatty Acid	
amodus.			

Flexible Gloss Wood La	cauer	"Aquarell" Colors
Nitrocellulose, Wet (1/4 sec.) 14 g.	Pigments
Ester Gum Solution (4# cut	20 g.	
Blown Castor Oil	4 g.	White:
Dibutyl Phthalate	$\hat{3}$ $\hat{\mathbf{g}}$.	Whiting Finest, or China Clay.
Ethyl Acetate	10 g.	Pale Yellow:
Butyl Acetate	10 g.	Pale Yellow Lake, or Yellow
Butyl Alcohol	7 g.	Lake, Blended.
Toluol	32 g.	Yellow:
	o . 6.	Yellow Lake, Martius Yellow.
Ethyl Cellulose Wood La	cquer	Ochre.
Ethyl Cellulose (Low Vis-	-	Pale Orange:
cosity)	8 g.	Orange Lake, Blended to get
Dewaxed Damar Gum Solu-		Lighter Colors.
tion (4# cut)	12 g.	Orange:
Dibutyl Phthalato	2 g.	Orange Lake.
Alcohol	10 g.	
Toluol	58 g.	Rosa (Pink):
"Cellosolve" Acetate	10 g.	Alizarin Lake, or "Echt-Rot"
	B.	(Genuine-Red), Blended to Ob-
		tain Lighter Colors.
Flat Wood Lacque		Red:
Nitrocellulose, Wet (1/4 sec.) 12 g.	Alizarin Red, Martius Red.
Dewaxed Damar Gum Solu-		Pale Brown:
tion (4 lb. cut)	10 g.	Terra di Siena, Blended
Ester Gum Solution		Brown:
(4 lb. cut)	10 g.	
Blown Castor Oil	2 g.	Caput Mortuum (Iron Oxide).
Dibutyl Phthalate	1 g.	Dark Brown:
Halowax No. 1014	5 g.	Umbra, or Cassel Brown.
Ethyl Acetate	5 g.	Violet:
Butyl Acetate	15 g.	Brilliant Violet Lake.
Butyl Alcohol	7 o.	Pale Blue:
Toluol	25 g.	Blue Violet Lake, Blended.
Xylol	8 g.	Blue:
Flexible Barrel (Inside)	Tuntin m	Blue Lake.
		Dark Blue:
a. Gilsonite Asphalt	50 g.	Dark Blue Lake.
Benzol	50 cc.	
b. Caoutchoue, Crude	5 g.	Pale Green:
Benzol	50 cc.	Green Lake, Blended.
Prepare a in an iron kettle	with stir-	Green:
rer, if necessary, heat.		Green Lake.
Prepare b soaking cold f	or several	Gray:
Prepare b soaking cold f days. Mix the two viscous	solutions,	Black Lake, Blended.
pouring b into a, stirring vig	orously.	Black:
Apply repeatedly, allowing	each layer	Black Lakes.
to dry well.	·	The blending, to get paler shades, is
		done by mixing the lake or pigment with
Inside Coating for Wood	Barrels	white chalk.
(Yellow Wax	40 g.	
		Manufacture of "Aquarell" Colors
Colophony	200 g.	
b. Iron Oxide	40 g.	(Water soluble, applied with brush)
c. Gypsum (Molding)	10 g.	Solution for binding of the pigments
Melt up a, then stir in b,	finally c.	in the color-paste:
Apply liquid, hot mixture wit	h a brush.	Formula No. 1
		04.11.
Lacquer for Barrels	4	Water, Distilled 51.9 g.
Rosin	22 lb.	
Turpentine, Thick	4 lb.	
Turpentine, Thick	4 lb.	Beef-Gall, Prepared 4 g.
Alcohol	12 lb.	Moldex or Other Preservative 0.1 g.
		B

Dissolve gum powder in cold water, stir, then heat to get complete solution. Add preservative, then glycerin, glucose solution, beef-gall. Filter, when cooled, through a percolator-cloth. (See No. 2)

No. 2

Dextrin, White	40 g.
Water, Soft or Distilled	41.8 g.
Borax, Crystallized	2 g.
Glycerin (28° Bé.)	6 g
Glucose Solution (1:1)	10 g.
Moldex or Other Preservati	ive 0.2 g.
Make dextrin paste in cold	water, then
earm to get clear solution, ad	ld preserva-
ive and horay then glucoses	alution and

glycerin.

Add the amount of water lost by evaporation (also in No. 1).

Alkali and Acid Resisting Paints

Formula No. 1	
Chlorinated Rubber	18 lb.
Toluol	43 lb.
Turpentino	9 lb.
Tetralin	4 lb
Wood Oil Stand Oil	9 lb.
Red Pigment	17 lb.
Amyl Acetate	1 lb.
No. 2	
Chlorinated Rubber	18 lb.
Toluol	45 lb.
Gutta-Percha Resm	11 lb.
Wood Oil Stand Oil	2 lb.
Amyl Acetate	6 lb.
Tetralin	5 lb.
Paint Graphite	11 lb.
Carbon Black	1 lb.

Fireproof Paints (for Wood)

Barium Sulphate	25 oz.
Zinc White	1 oz.
Water	20 oz.
Waterglass	25 oz.

Heat Sensitive Paints

Certain chemicals in form of paints can be employed for the detection, or determination, of temperature fluctuations of a surface. Thus, the double iodide of silver and mercury, which is yellow at ordinary atmospheric temperatures, is colored dark orange on heating, being brick red at a temperature of 70 to 80° C. The double iodide of copper and mercury is bright red at ordinary temperatures, turning chocolate brown at 70° C, and black at 100° C. If the heating of the paint films is not ex-

tended too far, the original color of the paint returns on being cooled back to ordinary atmospheric temperatures. A process recently patented in France employes a mixture of two substances, which react upon each other at elevated temperatures only, lead sulphide and barium superoude. In a suitable carrier this mixture is black at ordinary temperatures, turning gray on heating. This change is due to the formation of lead sulphate in the mixture.

Lime Resistant Paint

Complete protection against corrosion by hot line water and neetylene readines is obtained by a paint continuing 16 per cent chlorinated rubber, 44 per cent xylene, 35 per cent hthopone, and 5 per cent tritolylphosphate.

Luminous Paint Swiss Patent 172,076

This I did no 115,01	U	
Sandarae	36	g.
Rosin	18	
Paruffin		g.
Alcohol	35	
Petrolenm Ether	10	g.
Tricresyl Phosphate		g.
Benzom, Gum	2	g.
More middle and the amount of		12

Mox with gentle warming until dissolved. Dehydrate with quick lime and filter.

65 grams of above are mixed with:

Mildew Preventatives for Paint

Strontrum Sulphide

The addition of any of the following per 600 pounds of paint is advisable:

Mercuric Chloride	1 lh.
Sodium Silico Fluoride	6 lb.
Ammoniated Mercury	2 lb.

Non-Caking Pigments

Pigments are prevented from caking and are more readily dispersed in either oil or water if they are suspended in a dilute dispersion in water of diglyeol stearate or glyceryl monostearate and then dired. A film of waxy material is formed around each pigment particle. This film is both oil soluble and water dispersible.

Marble Effect Dipping Paint

Beautiful, marble like effects are obtained by dipping objects into many-colored paints floating upon the surface

of water. In order to float on water, the paints used have to weigh less than 8.33 pounds per gallon. Assuming that a varnish is used which weighs 7 pounds per gallon, the following table gives the number of pounds of pigment which, will yield a paint of sufficiently low weight to float on water, and have good hiding.

Chrome Yellow	1.25
Chrome Green	1.00
Prussian Blue	0.50
Para Red	0.50
Aluminum Bronze Powder	1.50
Gold Bronze Powder	1.50
Carbon Black (High Strength)	0.50

The procedure is important. Select a container which is wide enough and deep enough to hold the largest object to be dipped. Fill the container with water at room temperature. By means of a rod or dropper place a few drops of a colored paint here and there on the surface of the water. Near these drops or upon them place drops of a contrasting colored paint. Three, four or even five different colors may be used, but an excess of paint should be avoided. The colors will spread about, innighing with each other. They may also be blown gently. Hold the object to be decorated in such fashion that the entire outside surface is exposed. Immerse it slowly into the colors and into the water, turning it a bit at the same time. Blow the remaining colors aside in order to withdraw the object without having it traverse the colors again. The designs produced in this manner will always be different from each other, and are almost impossible to reproduce by hand painting.

Oiticica Oil Emulsion Paint U. S. Patent 1,998,845

. ,
120 oz.
6 oz.
2 oz.

Heat to 250° C, and then reduce to 200° C, and add

oo" C. ana aaa	
Potassium Silicate	13 oz.
Milk of Lime	16 oz.
Water	sufficient
Agitate violently until	Lool.

Paint Perfume

Vanillin is dissolved in turpentine or linseed oil. One part of vanillin is used to 2000 parts of paint to cover objectionable odors.

Plastic Paints

Zinc White or Lithopone	18.15	oz.
Water	7.5	oz.
Hide Glue	0.68	oz.
Linseed Oil, Pale Boiled	3.8	oz.
Rosin (WW or WG)	3.6	oz.
Benzol	3.8	oz.
Zinc Sulphate	0.12	oz.
If a hard dry product is	wishe	d, add

If a hard dry product is wished, add gypsum. Treat with water until pasty.

Synthetic Resin Enamel Paints Formula No. 1

Thin Stand	(White Seal) Oil	400 lb. 180 lb. 100 lb.
Turpentine		100 lb

Pug well and grind four times, then add:

China Wood Oil Varnish, con-

tauning 25 per cent Syn thetic Resin, equivalent to		3 lb	
Thick Stand Oil	40	lb)	
Turpentine		l lb	
Cobalt Linoleute (Liquid)	20	lb)	
This enamel dries in from	15	to	18

hours.

No. 2	
Titanium Oxide	300 lb.
Zinc Oxide	300 lb.
Thin Stand Oil	180 lb.
Synthetic Varnish	250 lb.
White Spirit	100 lb.
Cobalt Linoleate	10 lb.

No. 1

No. 3	
Zinc Oxide	300 lb.
Titanının Oxide	300 lb.
Thin Stand Oil	280 lb.
Synthetic Varnish	150 lb.
White Spirit	100 lb.

Synthetic Resin for Paints Canadian Patent 348,347

Castor oil 500 and drying oils 500 parts by weight are mixed and distilled until the residue of polymeric esters is approximately 85% of the original mixture. The retort is cooled below 290° and 800 parts of glycerol is gradually introduced. The mixture is heated for a short time well above the boiling point of water but below the boiling point of glycerol, and then 1200 parts of phthalic anhydride is gradually added, the temperature being maintained about midway between the boiling point of phthalic anhydride and that of water. When the mixture is clear and homogeneous it is run into cooling pans or into mixing tanks to be thinned with solvents.

Tar and Asphalt Paints Formula No. 1

Pine tar 120 l., rubber (small pieces) 1300 g., gutta-percha (small pieces) 1600 g., shellac 2700 g., copal varnish 4.5 l. When the varnish has been incorporated 45 l. of linseed oil heated separately to nearly the same temperature are added slowly.

No. 2

Asphalt 40 g., fossil resin 10 g., heat-thickened linseed oil 8 g., liquid driers 20 g., turpentine 60-70 g.

Paint for Marking Wood Boxes, Barrels, etc. Formula No. 1

Gum Arabic	3			10 g.	
Soda Ash				1 g.	
Glycerin				1 g.	
Water				40 g.	
Lampblack	or	pigment,	as	much a	1.5

Lampblack or pigment, as much as needed.

No. 2

Waterproof: 60 g. Shelhac, Ruby 60 g. Borax 60 g. Water 750 g. Dissolve boiling, and add: 60 g.

Gum Arabic 60 g. Pigment or Lampblack, as much as needed.

Cement Water Paint German Patent 575,895

Silica	40 kg.	
Pyrolusite	5 kg.	
Whiting	40 kg.	
Cement	15 kg.	
Grind very finely	and mix into the	e
following solution:		
Casein	50 kg.	
Borax	30 kg.	
Water	150 kg.	
Rosin Emulsion	20 kg.	

Wool Fat Emulsion Paints German Patent 612,715

Ammonium salts of high molecular fatty acids derived from drying or semi-drying oils have been claimed to be exceptionally valuable emulsifying agents for paint compositions incorporating both wool fat and non-water-soluble ingredients, such as resins and drying oils. Not only are the resulting coatings far more water-resistant than those of ordinary wool fat coatings, but the employment of an aqueous medium obviates

some of the drawbacks of solution in organic solvents. The process can be illustrated with reference to an enulsion of crude wool fat, refined tung oil and rosin, which are melted up in the respective proportions of 360:40:250, the melt being incorporated with 43 parts of ammonium solution, 100 parts alcohol and 1207 parts water and the resulting emulsion agitated till cold. The product at this stage, a viscous, yellowish-white emulsion, may be directly employed as a paint. An example of a quick-drying paint comprises 1000 parts emulsion, S0 parts chrome oxide, 150 parts itianium white and 15 parts of a 33 per cent solution of a cobalt-lead-manganese drier. Such a paint is stated to reach surface dryness within two hours after brushing on any type of surface, and admirably resists the action of a condensed steam-laden atmosphere.

Specialty Paints

French Patents 44,177 and 756,535

Under-Water Paint:	
Water	500 kg.
Tar	300 kg.
Caoutchoue Solution	200 kg.
Rosin	200 kg.
Benzene	100 kg.
Alum	2 kg.
"Very Brilliant" Paint:	
Alum	12 g.
Aluminum Bronze	5 g.
Salt	30 g.
Sugar	5 g.
"Fatty" Lime	50 g.
Water	400 g.
Oil	400 g.
Rosin	200 g.
Benzene	150 g.
Mica Powder	20 g.
Milk Whey	100 g.
Caoutchouc Solution	200 g.
Liquid Drier	150 g.
Pigments	10-15 g.

Paint and Varnish Remover Formula No. 1

15 lb.

14 lb.

Amyl Acetate

Paraffin Wax

Acetone .	14	lb.
Benzol	11	lb.
Methanol	12	lb.
Paraffin Wax	21/2	lb.
No. 2		
Whiting	21	lb.
Acctone	21	lb.
Denatured Alcohol	21	lb.
Benzol	23	lb.

No. 3

A low priced and effective remover tay be made up as follows:

of me made up do rononer		
Ethyl Acetate	30	oz.
Benzol	40	oz.
Methanol	27.5	oz.
Paraffin Wax	2	oz.
Methyl Salicylate	0.5	oz.

The parafin is melted and poured into he benzol. The other solvents are mixed nd then the benzol wax solution added o same while mixing vigorously.

Removing Plastic Paint

Mix one pound sal soda and two ounds hydrated lime and one-fourth of pound of table salt. Add enough rater to this mixture to produce a fairly cavy paste. Apply the paste with a beer brush, and leave it on until the old aaterial is softened, when it may be craped off. If the paste material should econe nearly dry before the old material is soft enough to be easily scraped off, apply the paste material again, but laways be sure you do not get this austic paste on the woodwork or floors, so it would injure them. When all the old material has been scraped off, wash he surface and rinse it until it is perfectly clean, and allow it to become dry sefore applying the first coat of paint.

Finish Remover

U. S. Patent 1,974,744

35 oz.
15 oz.
10 oz.
10 oz.
10 oz.
20 oz.
4 oz.

Varnish Remover, Liquid

Methanol	30 gal.
Phenol (90%)	5 gal.
Light Coal Tar Oil	65 gal,

Varnish Remover, Paste

Crude Vaseline	50 gal.
Phenol (90%)	45 gal.
Fusel Oil	20 gal. 80 lb.
Wood Flour	80 lb.

Shellac Finish

Shellac	250	g.
Dragon's Blood	50	ğ.
Alcohol	750	ğ.

Mix until dissolved, while warming on water bath.

Copal (Powdered and Exposed to Air for a Few Weeks)

Alcohol 250 g.

Dissolve by mixing on water bath and

then add:
Chalk, Precipitated 180 g.

Then mix with first solution.

Flat Indoor Shellac Lacquer

ritt	rngoor	Shellac	racquer	
Copal			131/2	oz.
Alcohol			131/2	oz.
Shellac T	l.N.		7	oz.
Alcohol			18	oz.
Bone Oil	l		3	υZ.

Flat Outdoor Shellac Lacquer

Shellac, Orange T.N.	50	oz.
Alcohol	200	oz.
Bone Oil	5	oz.
Oxalic Acid	1/2	oz.

Finishing Shellac Lacquer

Shellac, White R	efined 100	oz.
Alcohol	125	oz.
Butyl Alcohol	4	oz.
Bone Oil	1	oz.

Brushing Finishing Shellac Lacquer

Copai	21/2	OZ.
Alcohol	21/2	oz.
Sandarac	1/2	oz.
Alcohol	1	oz.
Shellac, T.N.	2.2	oz.
Alcohol	3.3	oz.

Acaroid Red, Alcoholic
(1:1)

Acaroid Yellow, Alcoholic

Shellac Floor Finish

280 g.	
50 g.	
1 l.	
	280 g. 80 g. 50 g. 1 l.

Stir altogether, let stand over night.

Floor Refreshener

5 lb. Shellac "Cut" Denatured Alcohol					gal. gal.	
This	mixture	is	applied	with	8	mop

This mixture is applied with a mop. The alcohol cleans and at the same time there is left a thin film of shellac which adds lustre to the floor.

Shellac Polish

Lac, Button	18 oz.
Alcohol	72 oz.
Shellac, T.N.	9 oz.
Sandarac	4 oz.
Benzoin, Gum	4 oz.
Turpentine, Venice	5 oz.

Water Shellacs

Water offenace	
1. Bleached "Pig-Tail" Sl	ellac
Water	645 g.
Borax	55 g.
"Pig-tail" Shellac, Ground,	
20% Water	300 g.
2. Bleached Shellac Power	der
Water	705 g.
Borax	55 g.
Shellac Powder, Dry	240 g.
3. Ruby and Orange She	llac
Water	700 g.
Borax	50 g.
Ruby or Orange Shellac	
(Free of Rosin and Wax)	250 g.

Solution in above formula is hastened by warming and stirring.

Water Resistant Shellac

Add 2-3% of urea or thiourea to solution of shellae in alcohol.

Bleaching Shellac

Lac may be bleached by dissolving it in 2.5% sodium carbonate solution at and, after filtration and cooling to air temperature, adding a solution prepared by passing chlorine into a solu-tion containing 12.5% of caustic soda and 2.5% of sodium carbonate. latter should contain 6-8% of available chlorine and, if of pH 10-10.5, does not require storing in a cool place. The amount of such a solution necessary for bleaching indicates a chlorine requirement of 10-14% on the weight of lac, and a yield of 93-95% is obtained. The bleached lac may be recovered by the slow addition, with stirring, of 1:20 sulphuric acid, the precipitate being then collected, washed, and dried in vacuo over sulphuric acid. The product is freely soluble in cold 97% alcohol, and the solubility does not alter on prolonged storage in air. The bleached material contains 2.3-3.1% of moisture, 0.98-3.52% chlorine and has a saponification value 236.0-256.7, acid value 70-68-83-52, and iodine value 3.9-5.0.

Substitute Shellac Solutions

The substitutes for shellac solutions are of three types:

- 1. Substitute for wax-free shellac solution.
- 2. Substitute for white shellac solu-
- 3. Substitute for orange shellac solution.

The base for all three is the same, namely a solution of a cheaper alcoholosoluble resin in completely denatured alcohol. At the present time a soft Manula gum is used, and a 6-lb. cut represents the maximum concentration normally made. To prevent loss by evaporation, as well as to avoid the hazard of volatile alcohol vapors, a closed tumbler is used, in which is placed one gallon of alcohol for every six pounds of the Manula gum. When solution is complete, the tumbler is emptied and the solution allowed to settle. The clear supernatuat solution represents a substitute for wax-free shellae solution.

White and orange shellae solutions contain a cloud of suspended wax which is inherent in the material and insoluble in alcohol. To duplicate the waxy appearance a preparation of carnauba wax may be employed. A quick and safe method of preparing the wax is as follows:

Imitation Shellac "Cloud"

Dissolve 5 lb. of carnauba wax in onehalf gallon of blown castor oil. Since carnauba wax melts at 84-86° steam-jacketed kettle may be used. If a direct fire is used, the flame must be extinguished before proceeding further with the formula. Add slowly and with constant stirring one-half gallon of turpentine, followed by one balf gallon of denatured alcohol. A soft yellowish-white paste will form. This paste, added to a solution of 95 lb. of Manila gum in 15 gal. of alcohol, represents a 6 lb. cut in which the wax constitutes 5% of the total solids. Less paste may be used, but not more. The castor oil serves not only as a solvent for the wax, but also as a plasticizer.

The waxed product is a substitute for white shellac. It may be colored by means of an orange alcohol-soluble anilone dye, thus forming a substitute for orange shellac.

Shellac Substitute

U. S. Patent 1.942,413

Batu (Galla-Galla) Gum 18-20 oz. Rosin 10-20 oz. Heat to 260° C. Add: Calcium Oxide

1-4 oz.

Heat to 320° C. and stir till dissolved. Cool and "cut" with varnish solvents to give a shellac substitute solution.

Oiticica Varnish

An oiticica oil varnish cooked under the same conditions as a similar tung oil varnish is lower in viscosity, which is an advantage. If the temperature is taken over 250° C. frothing occurs and this has to be carefully watched.

By blowing oiticiea oil for 30 minutes at 220° C, a thick light-colored oil is formed which will be comparable with blown linseed oil. Oiticiea oil varnishes have a less characteristic odor and are less noticeable in closed spaces.

To establish the technical value of officien oil, tests have been made with varnishes with a natural or artificial resin base and mixtures on the one hand of tung oil and linseed oil, and on the other of oiticien oil and tung oil, the latter being in the ratio of one part to two respectively. Heating is done at 315° C. and maintained until the mixture has the correct body.

Ester Gum Varnishes Formula No. 1

Ester Gum	100 lb.
Tung Oil	198 lb.
Linseed Oil Heated for	
2 Hours	36 lb.
Solvent Naphtha	81 lb.
White Spirit	250 lb.

Driers are added in the proportion of 0.5% lead and 0.035% cobalt. This gives a varnish which becomes tacky in 45 minutes and dues in about 3 hours. The film is resistant to cold and boiling water. The film is not resistant to combustion gases. The Gardner-Holt viscosity is D and the color 11.

No. 2

Ester Gum	100 lb.
Oiticica Oil	156 lb.
Tung Oil	78 lb.
Solvent Naphtha	84 lb.
White Spirit	250 lb.

Driers are used in the same proportion, i.e., 0.5% lead and 0.035% cobalt. This varnish becomes tacky in 2 hours and perfectly dry in about 8 hours. The film is resistant to cold water but not to boiling water when it whitens but becomes transparent again.

Pharmaceutical Cellulose Varnish French Patent 777,999

A varnish containing, e.g., benzylcellulose 5-18 g., benzine 18-40 g., tolueno or xylene 25-45 g. and butyl acctate 20-35 g., or benzylcellulose 2-12 g., benzine 50-80 g., and ether 25-80 g., used for pharmaccutreal or toilet purposes, is contained in a collapsible tube and used as required.

Electrically Conducting Varnish Formula No. 1

Aluminum Bronze Powder	240 g.
Synthetic Resin Varnish	1 l.
No. 2 Copper Bronze Powder Lacquer	120 g. 1 l.

Cold "Cut" Synthetic Resin Varnish
Rezyl No. 14 10 g.
Methanol 50 cc.
Toluol 50 cc.

Allow to stand over night and then stir.

Leather Roller Varnish

reguler v	oner varnisn	
Venetian Red	4	lb.
Ground Blue	5	lb.
Vinegar	15	
Glycerin	75	
Glucose	150	
Oil of Cloves	25	
Methyl Saheylate	25	cc.

Mu Oil Varnish

Mu Oil 200 oz. Modified Phenolic Resin 100 oz.

Heat with stirring to 570° F.; keep at this temperature for 6 minutes; cool to 350° F. and dilute with 250 oz. petroleum spirits and add 5½ oz. lead naphthenate and 0.1 oz. cobalt naphthenate.

Mopping and Wiping Varnish

Because varnishes of this type leave a very thm film, it is essential that they be made of tough and durable ingredients. The average floor or furniture varnish, if thuned to wiping consistency, is unsuitable. A high grade product consists of a blend of 3 or 4 pints of the following varnish a with 1 pint of varnish b.

a. Bakelite XR-4070 100 lb.
China Wood Oil 16 gal.
Body for 1 hour at 450° F. Reduce
with 20 gal. mineral spirits, 5 gal.

xylol, 3 gal. dipentene, 2 gal. high boiling hydrogenated naphtha and 3 gal, gum spirits of turpentine.

b. Substitute Bakelite BR 820 in place of XR-4070 in Formula a, and body with the wood oil at 400° instead of

Driers are unnecessary.

High Gloss Transparent Printing Varnish

British Patent 426,7	53
Ester Gum	120 oz.
Tung Oil	40 oz.
Linseed Oil (Half Boiled)	40 oz,
Mineral Spirits	6 oz.
Cobalt Linoleate	5 oz.

The above may be colored with a suitable amount of rhodamine base dissolved in olein, Berlin blue, alizarin madder lake, milori blue or Sudan yellow.

Wrinkle Finish Varnish U. S. Patent 1,934,034

100 oz. Tung Oil 5-10 oz. Rosin Heat for 2 to 8 hours at 177-290° C.

Cool and dissolve in an equal weight of high-flash naphtha.

Limed Rosin

The apparatus and procedure vary somewhat, but the following is usual practice: One hundred and twenty five pounds of resin are melted in a cylindrical flat-bottomed copper vessel 36 inches in height and from 30 to 36 inches in diameter. The vessel has a loose cover provided with a small chimney and an opening for the stirring rod. It is mounted on an iron truck, the platform of which is about 2 inches from the floor. The truck is then wheeled to a position under a chimney and over a furnace, which is located beneath the floor. The resin completely melts in about a half hour. It is at this point that the use of lime enters.

Lime is added, gradually, to the melted resin with the temperature at about 350° F. Theoretically, about 13.6 pounds of hydrated lime would be required, but it is inadvisable to completely neutralize the resin. In actual practice 8 to 10 pounds of hydrated lime are used. This reduces the acidity of the resin from about 160 to 65. After stirring and heating for a short while, the treatment with lime is completed.

Wood Filler

Shellac (if for Transparent Wood Filler Use Blenched Shellac) lb. gal. lb. Methylated Spirits 20 Barytes Silica 10 lb. Raw Linseed Oil 1/4 gal. Dissolve the shellar in the methylated

spirits and add the Imseed oil. Mix the burytes and silica together dry, and stir into the shellae varmsh. Grind to a smooth paste and adjust the consistency with additional barytes and silien mixture or shellac varmish. Store in airtight containers.

Filler-Undercoat for Shellac

Mixing powdered bornele acid, 5 g., with each ounce of shellar to be used as an undercoat on wood causes the shellne to dry very hard so that it serves as a filler as well as an undercoat.

Porous Wood Scaler

One handred thirty-five pounds of 400mesh Silica, 65 lb. Bentonite, 10 gal. of Congo Copal Varnush, 2½ gal. Pontinnak Gum Varnish, 2½ gal. Nevindene Solu-tion, 10 gal. Light Nuphtha, 5 gal. Lacquer Thuner, 12 gal. Concentrated Cobalt Drier. Nevindene solution is (basis) 6 lb, of Nevindene resin cut cold in 1 gal. of mineral spirits.

The protective covering should be a coat of aluminum paint and advisedly two conts of regular oil-type house paint. The Aluminum Primer recommended is: 7212 gal. of an 80-gal. Tung/Ester Varnish, 10 gal. Boiled Lanseed Oil, 7 gal. Mineral Spirits, ½ gal. Lead-Manganese Concentrated Drier, 135 lb. Paste Alumimm (or powder).

Non Penetrating Plaster Sealer

45 lb.

Pigment.

Velnele	55	lb.
Pigment:		
Titanium-Calcium Pigment	62	lb.
Metronite	37	lb.
Aluminum Stearate	1	lb.
Vehicle:		
Bodied Linseed Oil	50	lb.
Mineral Spirits	45	lb.
Liquid Drier	5	lb.

Wood Filler for Ground Polishing German Patent 607,521

Shellac Wax	10	oz.
Carnauba Wax	5	UZ.

	Pumice Meal			100	oz.
	Sandarac			100	oz.
	Blown Castor	Oil		10	oz.
,	Melt together	until	uniform	and	pow

vder after cooling.

American	Walnut	Graining	Color
Ivory Blac	k	2	oz.
Van Dyke I		4	UZ.
Burnt Umb	er	2	oz.
Bolted Wh	iting	1	OZ.
Water	•		1/2 gal.

Imitating Old Copper Finish After application of priming coat use White Lead lb. Chrome Yellow, Medium 12 oz. Venctian Red 11/2 lb. Burnt Umber oz. 41/4 lb. Linseed Oil Turpentine 41/4 lb. to suit Drier

After applying above paint, allow to dry and use a coating of copper bronze powder thinned with equal parts of spar varnish and turpentine. When this coat is dry apply a glaze made from chrome green, medium, thinned with equal parts of raw linseed oil and turpentine plus a small amount of drier. While the glaze is still damp wipe it here and there to produce a mottled effect.

Liquid Oil Graining Color

Raw Linseed Oil		gal.
Turpentine	3/6	gal.
Drier, Liquid		pt.
Beeswax, Yellow (Shavings)	₹3	0 Z.
Warm together and mix until	cl	ear.

Wood Stain

U. S. Patent 1,977,345

Dye, Water Soluble 4 oz. Diethylene Glycol Ethyl Ether 5 oz. 80 oz. Ethylene Glycol Methyl Ether 15 oz.

Wood Stain

U. S. Patent 2,000,121 Diethylene Glycol Mono

netnylene Glycol	MIOHO-		
ethyl Ether		1	oz.
Lethyl Alcohol		9	oz.
Coluol		6	oz.

This composition may be utilized with from 2 to 2½ oz. of the particular dye to 1 gal. of the composite solvent. The amount of dye utilized depends on the particular dye itself and its degree of

concentration, and the depth of color required in the particular stain. Further, the strength of the dye stain may be varied by the amount of diethylene glycol mono ethyl ether utilized.

In making up these compositions, the aniline dye or stain, such as the nigrosines, may be allowed to stand with the diethylene glycol mono-ethyl ether until the dye dissolves, after which the other ingredients may be added.

Coloring of Light Wood to Imitate Ebony

A vacuum process is essential for good impregnation of wood with coloring substances. Aqueous solutions are preferable where possible on grounds of low price, high vapor pressure (which assists impregnation) etc. Woods for this ebonizing process, in order of suitability are: apple, pear, hazel, maple, beech and birch. The following are recipes for ebonizing:

Formula No. 1

Gall-nut solution containing a few drops of ammonium vanadate solution.

No. 2

3.60 kilograms andline hydrochloride, 1.80 kilograms potassium chlorate, 40 liters water, 0.250 liter hydrochloric acid, 4.20 grams ammonium vanadate.

Four kilograms carbon black, 18 liters shellae Japan lacquer, 18 liters turpentine.

No. 4

1,200 kilograms carnauba wax, 3 kilograms ceresm, 30 grams oil-soluble nigrosine, 10 liters turpentine.

Auto Top Dressing

Orange Shellac Denatured Alcohol gal. Castor Oil 1/2 oz. If a black finish is desired add nigrosine to give the desired color.

Butter Taint Prevention Coating Tubs are coated with following: Prime Lactic Casein 50 oz. Borax 7.5 oz.

Water Stir and warm gently until smooth.

> Candy Glazes Formula No. 1

Sandarac 125 g. Benzoin, Sumatra

COMITTION, I			
Turpentine, Venice	10	Œ	Г
	740	e.	l
Alcohol No. 2	140	g.	
Benzoin, Sumatra	200		
Balsam, Peru	5		
Alcohol	800	g.	ĺ
No. 3		**	
Benzoin, Sumatra	150	g	١,
Shellac, Refined	50		١,
Vanillin	1	g.	
	800	p.	
Alcohol	500	ь.	
Protective Food Coating	p r		
			l
French Patent 780,762			l
Lactic Casein	100	g.	١,
Borax or Sodium Phosphate	16		١ '
Sodium Bicarbonate	32	g.	l
	34	g.	
Glycerin Distilled Water	820		١.
	8	g.	
Gelatin			ı
This may also be applied to a or tin foil for use on foods.			
Protective Coatings for Sauss	ıges,	ete.	
Formula No. 1			١
	5	g	1
Gelatin		g.	l
Salt	ī	g.	ı
Saltpeter No. 2	•	6.	
	Γ.	g.	1
Gelatin	1	g.	l
Glycerin	1	g.	l
No. 3			ı
Pentosan Resin		g.	1
Gelatin	1	g.	1
No. 4			ı
Aqueous Solution of Stahr, o Agar, or Gelatin, ½-2% For	r Ag mic	ar Acid.	
No. 5			
Tallow			١
No. 6			
Alum		g.	1
Olive Oil	1	cc.	1
Shellac	16	g.	1
Alashol		ec.	1

No. 9 Glue, Gelatin or Isinglass, boiled in a little vinegar.

No. 7

No. 8

Colophony, Shellac, Glycerin,

Alcohol

Paraffin

Colophony

Linseed Oil

or Wax

Whiting

65 cc.

35 g. 62.8 g.

2.2 g.

60 g.

40 g.

Laboratory Table Top Stain Solution No. 1

Potassium Permanganate 20 g. Copper Sulphate 20 g. Water 1 l.

Heat to about $60-70\,^{\circ}$ C, and apply to clean desk top, and follow immediately with solution No. 2.

Solution No. 2

Hydrochloric Acid	
(sp. gr. 1.2)	150 cc.
Ambue	150 cc.
Water	700 ec.

Heat to about 60-70° C. and apply over No. 1.

When the desk top is dry it may be rubbed with hissed oil in the usual manner.

Red Stamp Pad Ink

Fuchsin	1	OZ.
Glycerm	32	OZ.
"Lysol"	1/8	OZ.
Acetic Acid	1	oz.
Denutured Alcohol	1	oz.
Water	1	υZ

Protector for Polished Surfaces French Patent 778,389

Water	150 cc.
Linseed Oil	200 cc.
Alcohol	450 cc.
Sulphuric Acid	20 cc.
Shellac	30 g.

Coating for Old-Painted Surfaces Swiss Potent 173,070

Lithopone, as Pigment	optional
Benzohne	25-30 cc.
Polishing Lacquer	25 cc.
Trichlorethylene	25 cc.
SWISS Latent 115	,010

Preparation of Oil Pastes from Pigment-Water Pulp

The addition of linseed oil of acid value about 10 will cause the separation of water from a pulp of white lead-inwater. Agitation and friction are necessary in order to insure contact of the oil with the pigment and in order to express the maximum amount of water. With other pigments, particularly those whose affinity for oil is less striking than that of white lead, transfer from that the pigment is the striking than that of white lead, transfer from the striking than that of white lead, transfer from the striking than that of white lead, transfer from the striking than that of white lead, transfer from the striking that the strikin

accomplished by one or more of the following means:

- 1. High acid linseed oil.
- 2. Polymerized linseed oil.
- Linseed or China wood fatty acids.
- 4. Addition of various chemical agents.

As an example of method 4 (Patented), 15.5 parts of linseed oil (and value 7) or of other drying oil (and value gieater than 4) are added gradually at 82-88° C., with vigorous agitation, to a suspension of 100 parts of lithopone in 200 parts of water which also contains tri-sodium phosphate or other alkaline saponifying agent. water separates in the upper layer after 10 to 30 minutes' further agitation.

Strong Lead Oil for Black Paints

Varnish linseed oil is heated with continual stirring until at the end of an hour the temperature reaches 570° F. (==300° C.) and is held at this temperature for a further 3 to 4 hours, when the heat is closed down. Finely powdered white lead is then added slowly on a falling temperature, commencing at about 525° F. (=274° C.), in the proportion of 4½ lb. of white lead to every 100 lb. of oil, about 2 hours being occupied in adding the white lead. So far, it will be observed, the process will have occupied practically one working day. On the following day the oil is heated up again, care being taken to avoid local heating in the early stages until the whole mass becomes quite fluid. Hent is then increased until a temperature of 535° to 545° F. (==280° to 285° C.) is reached, at which the oil is held until the body required is attained. The purposes for which oil of this type is used demand as a rule that the product when cooked shall "string" very strongly when tested on glass. Gums or blacks with which it may be cooked afterwards are usually expected to "pill" between the finger and thumb.

Flatting Oil

Linseed Oil 15 lb. Solvent Naphtha or Turpentino 85 lb. Drier to suit

Add to the following lead paste in proportions of 21/2 gal. above oil to 100 lb. lead paste:

White Lead 92 lb. Linseed Oil 8 lb.

Black Iron Oxide Pigment Austrian Patent 141,130

Ferrous Sulphate 240 lb. 720 lb. Water Boil the above and while boiling add: Potassium Chlorate and then add. Sodium Carbonate 115 lb. `in Water 230 lb.

Various shades are obtained by varying composition of first solution, nature and amount of oxidizing agent and other reaction conditions.

Carmine Lake Pigment

Powder the best silver grav cochineal as finely as possible, and boil it for three hours in water. Filter the hot solution quickly through a thick linen cloth. Boil up the filtrate again, and add the substances needed to form the lake. Many such substances may be used, but only two can be thoroughly depended upon, and they should both be used together. These two are alum and tin salt, and if necessary, warmth may be given to the color by the cautious addition, drop by drop, of hydrochloric acid. The alum must be absolutely free from iron, or it will be impossible to get more than a very unsatisfactory product. The best proportions are:

Cochineal 20 lb. Water 500 lb. Alum (Iron Free) 2 lb. 2 lb. Tin Salt

The alum and tin salt are added at the boil, which is kept up till everything is dissolved. The clear solution is then exposed in shallow dishes covered with sheets of glass for several weeks in a very bright sunny place. By this time the dark-red liquor will have lost nearly all its color, and the carmine will have been deposited in the solid form, partly on the dish and partly on the surface of the hquid. It is separated by filtration, and carefully dried with blotting-paper. To get a fine and warm red it is absolutely indispensable that the dishes should get plenty of sun, so that the manufacture is impossible in any but the most favorable weather.

To get absolutely pure carmine, the product already described is dissolved in caustic ammonia. The solution is filtered, and the carmine is reprecipitated with acetic acid.

Satin White Pigment

Ninety pounds of quicklime are slaked in 27 gal. of boiling water. To this mixture 130 lb. of finely divided (260 mesh) aluminum sulphate are added quickly, and the mass is heated until it becomes almost solid. Thirty gallons of water are then added and the mixture agitated thoroughly. The last trace of any visible yellow color is neutralized by the addition of indanthrene blue mite form of a solution of 2.5 lb. of the commercial paste in 6 gal. of water. A very small amount of this solution is required if a good grade of lime and sulphate are used. The satin white is then filtered and dried.

Reflecting of Light by Colors

	Reflection
	Factor
Color of Paint	Per Cent
White (Gloss)	84
White (Flat)	82
White (Eggshell)	81
Ivory White	79
Cream	74
Aluminum (Made with Paste) 73
Ivory Tan	67
Light Green	62
Light Gray	59
Buff	55
Light Blue	52
Medium Green	49
Tan	48
Medium Blue	43
French Gray	32
· ·	

Printing in Several Colors British Patent 426,753

High-gloss color-printing is effected by printing the picture in black or other color in the usual way and over printing the picture entirely or partly with a transparent colored gloss overprint varnish. The varnish may be colored with oil soluble coloring matter or with highly glazing insoluble pigments or with both. In the last case, autotype prints having a double tone effect may be produced, the soluble color spreading out around each of the dots of the picture. The first print may be made with a normal black art printing ink. The varnish consists of clear resin ester 120, china wood oil 40, slightly boiled linseed oil 40, petroleum 6 and cobalt linoleate 6 parts. To 12 parts of varnish may be added 2 of rhodamine base in 2 of olein, 1 of Berlin blue or 1 of alizarin madder lake (I). Double tone effects may be produced by over-printing with a mixture of varnish 25, rhodamine base 0.5 in olein 0.5, and milori blue 1 parts, or with varnish 25, Sudan yellow 0.5 and (1) 1 part.

Dissolving Amber

The amber is powdered and heated under a reflux condenser with butyl alcohol containing a little hydrochloric acid for 6 to 8 hours.

Dustless Carbon Black

Formula No. 1		
arbon Black	200	g.
apropéhte Tar	24	g.

Water 50 cc.
Form pellets or briquettes and dry at 105° C. for 3 hours.

No. 2
Carbon Black 200 g.
Dextrin Solution (5%) 100 cc.
Treat as above.

Colloidal Preservative U. S. Patent 1,937,813

A transparent, solvent-resistant, antiseptic, colloidal mass is produced by condensing the gases evolved when gelatin 3 lb. or glue, etc., is heated with wood crossote 4 lb. at 160-250° C, for 2 hours.

Coloring Meerschaum Pipe Bowls

Beeswax	50 oz.
Olive Oil	50 oz.
Triethanolamine	15 oz.

The Meerschaum pipes are immersed in the above which is slowly heated to boiling and maintained at this temperature for 15 to 30 minutes. Pipes so treated will color very rapidly.

Blue Sheep Marking Pencil

Soapstone	28 1b.
Fine Gypsum	21 lb.
Chinese Blue	2 lb,
White Soan Powder	10 lb.

Mix all ingredients well together and make up with thin glue water into a stiff paste. They are then shaped like a thick pencil and dried.

Brewer's Glaze

Orange Shellac	25	oz.
Manila Copal	12	oz.
Acaroid Resin, Yellow or		
Red	8	oz.
Linoleic Acid	0.5	oz.
Alcohol	54.5	0 2.

Rubbing Compound

(For Paint, Lacquers, etc.)

1. Carnauba Wax	42 lb.
2. Beeswax	18 lb.
3. Ceresin	18 lb.
4. Varnolene	3 gal.
5. Water	3 gal.
6. Triethanola:nine	8 oz.
7. Stearic Acid	2 lb.
8. Tripoli	24 lb.
9. Pumice	15 lb.

Melt 1, 2, 3, 7 with 4. Heat 5 and 6 to 90° C., add to wax mixture and stir till emulsified. Then add 8 and 9 and stir till cool.

Peeled Wood Wall Paper U. S. Patent 1,945,686

The veneer is cut into strips of definite width which are dried, steeped in solution (1), dried, steeped in solution (2), dried, and finally backed with any kind off fibrous fabric. (1) comprises cellulose acetate 15, 14% solution of chrome alum 10, and water 70 oz., and (2) 25% glycerin 30, gelatin 25, and water 45 oz.

Double Strength Lead-Manganese Liquid Drier

Lead-Manganese Uversol 200 lb. No. 303 200 lb. Bodied Linseed Oil 73.5 lb. Pine Oil 9.0 lb.

 Pine Oil
 9.0 lb.

 Turpentine
 60.0 lb.

 Pine Tar Oil
 3.0 lb.

 Mineral Spirits
 254.5 lb.

Yield 75 gal.

This drier is double the strength of the preceding, containing 1.0% manganese and 11.0% lead as metals.

Procedure: Melt the drier quickly with the linseed oil at a temperature not exceeding 275° or 300° F. Remove from fire and reduce with the solvents.

Lead-Manganese Drier

Lead-Manganese Uversol

No. 303 100 lb. Mineral Spirits 500 lb.

Yield 85 gal.

This drier has an acid value =0. It contains 0.5% manganese and 5.5% lead as metals. One part of drier to twenty parts of oil will give a metallic content of 0.025% manganese and 0.275% lead.

COSMETICS AND DRUGS

COSMETICS	AND DRUGS	
Pine Needle Bathing Salt	Medical Bathing f	la lta
Formula No. 1	Carlsbad Well	
a. Salt 100 kg.	Sodium Sulphate	44 g.
b. Water, Containing 5%	Potassium Sulphate	2 g.
Uranin (Fluorescein-	Sodium Chloride	18 g.
Sodium) 2.5 kg.	Sodium Bicarbonate	36 g.
c. Sodium Carbonate, Anhy-		
drous 2.0 kg. d. Magnesium Carbonate 0.2 kg.	Friedrichshall	
c. Pine Needle Essence 2-3 kg.	Sodium Chloride	277
Mix a with b homogeneously, dry on a	Sodium Bromide	37.7 g. 0.3 g.
shelf and sift through a sieve, mix then	Potassum Chloride	5 g.
with c and d, in a drum, add c, mix	Calcium Chloride	19 g.
again thoroughly, fill into scaled cans.	Magnesium Chloride	37 g.
No. 2	Calcium Sulphate,	
Sodum Bicarbonate 10 g.	Precipitated	1 g.
Starch Powder 1 g.		
Tartaric Acid, Powdered 7.5 g.	Reichenhall	
Fluorescein or Uranin 0.1-0.2 g.	Potassium Chloride	6 g.
No. 3	Magnesium Chloride	6 g. 72 g.
Sodium Chloride 70 g.	Lithium Chloride	0.15 g.
Pine Needle Extract, Genuine 18 g.	Sodnim Chloride	14 g.
Ammonium Carbonate 10 g.	Sodmm Bromide	0.85 g.
Perfume (Pine-Needle) 2 g.	Magnesium Sulphato	7 g.
Ocean Bathing Salt		
***	Kreuznach	
(1000 g. per Bath)	Sodium Chloride	63 g.
Potassium Iodide 1 g. Potassium Bromide 0.55 g.	Potassium Chloride	75 g.
Lithium Carbonate 0.05 g.	Calcium Chloride	750 g.
Manganese Sulphate 0.01 g.	Magnesium Chloride Sodium Bromide	110 g.
Ferrous Sulphate 0.01 g.	Sodium Dromide	2 g.
Potassium Chloride 15 g.	***************************************	
Calcium Chloride 40 g.	Hallein Well	
Magnesium Sulphate 66.38 g.	Sodium Chloride	69.3 g.
Magnesium Chloride 96 g. Sodium Chloride 781 g.	Magnesium Chloride	27 g.
Sodium Chloride 781 g.	Sodium Bromide	0.42 g.
O 70 11: 01:14	Calcium Sulphate, Pre-	
Oxygen Bathing Salt	cipitated	10 g.
Formula No. 1	Sodium Sulphate	2.28 g.
Ammonium Carbonate, Dried 500 g.		
Hydrogen Peroxide (3%) 100 g.	Vichy	
Urea 5 g. No. 2	Lithium Carbonate	0.01 g.
Urea Hydrogen Peroxide 50-100 g.	Ferrous Sulphate	0.05 g.
Sodium Pyrophosphate 10 g.	Manganese Sulphate	0.01 g.
No. 3 (Tablets)	Sodium Chloride	1.73 g.
` ,	Sodium Sulphate	6.2 g.
Sodium Perborate 800 g. Starch 100 g.	Magnesium Sulphate Calcium Chloride	2.6 g. 6.0 g.
Ammonium Carbonate 100 g.	Sodium Bicarbonate	6.0 g. 83.4 g.
A3		P.

Mud Bath Salt		Steel (Iron) Bath	9
Ferrous Sulphate	900 g.	Formula No. 1	•
Calcium Sulphate, Pre-	5 ° 6.	Iron Tartrate	100 g.
cipitated	20 g.	Distilled Water	900 cc.
Magnesium Sulphate	20 g.	No. 2	000 00.
Sodium Sulphate	40 g.		20. 40
Ammonium Sulphate	20 g.	Iron Sulphate, Pure	30-60 g.
Optional, Dry Mud Earth.		Potassium Carbonate, Pure	120 g.
		No. 3	
"Saltrate Rodell"	,	Iron Sulphate	30 g.
Sodium Chloride, Powder	0.1 g.	Salt Sodium Bicarbonate	60 g. 20 g.
Magnesium Carbonate	0.5 g.	Bodium Bicarbonate	20 g.
Potassium Carbonate	0.1 g.		
Lithium Carbonate	0.05 g.	Sulphur Baths	
Calcium Sulphate, Powder	0.25 g.	Formula No. 1	
Borax, Powdered	10 g.	Potassium Sulphide	50 g.
Sodium Bicarbonate	30.5 g.	Eau de Cologne	50 g.
Ammonium Carbonate	52.5 g.	Distilled Water	950 cc.
Sodium Thiosulphate	2.5 g. 3 g.	No. 2	
Sodium Perborate	3 g.	Soft Soap	250 g.
		Glycerin	50 g.
Stimulating Bathing	Salt	Potassium Sulphide	25 g.
Sodium Chloride, Powder	950 g.	No. 3	
Sodium Bicarbonate	50 g.	Sodium Thiosulphate plu	ıs Acid
Thyme Oil	2 cc.	Bath-Water	
Bergamot Oil Terpenes	5 cc.	No. 4	
Orange Peel Terpenes	1 cc.	Sulphur Sublimed 5	60-100 g.
Bergamot Oil	1 cc. 1.5 cc.	Ammonium Carbonate 95	50-900 g.
Terpineol Methyl Naphthyl Ketone	0.5 cc.	Distilled Water Warm	650 cc.
Methyl Haphthyl Retollo	0.0 00.	b. Distilled Water, Warm Potassium Chromate,	000 00.
	D 41	Neutral	25-50 g.
Effervescent Tablets for	Batns	Mix a, dissolve b, mix bo	
Formula No. 1		several hours, until solid.	Press and
Sodium Bicarbonate	300 g.	grind; 120 g. used for a bath	•
Sodium Acid Sulphate	275 g.	No. 5	
Starch	25 g.	(Bain de la Parisier	ine)
No. 2	٠	Sodium Bicarbonate	870 g.
Saponin, Purified	2 g.	Magnesium Carbonate	10 g.
Starch Sodium Bicarbonate	25 g. 90 g.	Sulphur Flowers, Ground	
Tartarie Acid	70 g.	Sulphur Flowers, Ground Sulphur, Precipitated	20 g.
The stability can be in		Selenic Acid	0.1 g.
pressing the bicarbonate and	l acid sepa-		
rately.	Dollar	Carbon Dioxide Bat	hs
		Formula No. 1	
	*** ***	Ammonium Carbonate	35 g.
Effervescent Tablets with	Wetting	Sodium Bicarbonate	20 g.
Agents		Tartaric Acid	30 g.
(Slow Development of Carb	on Dioxide)	Sodium Perborate	10 g.
Formula No. 1		Sodium Thiosulphate	3 g. 2 g.
Starch	10 g.	Disodium Phosphate	4 g.
Sodium Lauryl Sulphonate) 10 g.	No. 2	40 -
Sodium Bicarbonate	46 g.	Sodium Bicarbonate	42 g.
Tartaric Acid	34 g.	Sodium Acid Sulphate	21 g. 5 g.
No. 2		Starch Sodium Chloride, Powder	30 g.
Sodium Bicarbonate	57 g.	No. 3	0. B.
Tartaric Acid	38 g.	1	95 ~
Saponin, Purified	5 g.	Ammonium Carbonate	25 g. 20 g.
Stearin, Powder	5 g.	Sodium Bicarbonate	20 g.

COSME	1105	AND DRUGS	- 65
Tartaric Acid 25	~	ditional untar until accident	
Sodium Perborate 10		ditional water until previous absorbed.	amount is
Rice Starch 20	g.	No. 5	
Manganese Nitrate			10 -
710 1		White Beeswax	12 g.
Mix all components—except the per	the	White Petroleum Jelly Peach Kernel Oil	12 g.
rate—dry and perfume, then add	the	Rose Water	50 g. 25 g.
perborate. Press in tablets.	- 1	Borax	
	- 1	Perfume	1 g. to suit
M ud Bath	ı	Terrumo	to buil
Ferrous Sulphate, Crude 900 g	c.		
Engom Sulta 20 o	œ l	Greaseless Cold Creat	m
Glauber's Salts 40 g	g.		16 oz.
Ammonium Sulphate 20 g	g.		18 oz.
Gypsum, Crude 20 g	g.		12 oz.
Clay, Dark 50 g	g.		2 oz.
			4 07.
That Dath Donalas (as Mallata) w			54 oz.
Foot-Bath Powders (or Tablets) w	with	Perfumo	.75 oz.
Perborato	- 1		
Formula	. 1		
No. 1 No. 2	1	Cold Cream	
Sodium Perborate 170 g. 180 g	g.	1. Diglycol Stearate 14	lb.
Boric Acid, Powder 70 g. 60 g	g.	O Dometto Way 0	
Borax, Powder 50 g. —	1	3. Mineral Oil 33	¼ gal.
Sodium Acid Car-		4. Petrolatum (White) 6	lb.
bonate 250 g. 200 g	g.	5. Water 6	gal.
Perfume 5-10 g	- 1	6. Perfume Oil 5	1/2 fl. oz.
Tablet or powder doses for each b	bath	Method of manufacture:	
should weigh 10-20 g.		a. Melt Nos. 1, 2, 3 and 4	nt 170° F.
	- 1	b. Heat 5 to 180° F.	
Cold Creams		c. Add b to a while mixir	ig. Allow
		mixer to run until bate	h is com-
Formula No. 1	j	pletely emulsified,	
Cetyl Alcohol 10 g		d. Allow batch to cool to 12	
Paraffin, Liquid 10 g	g.	add 6 and mix at low spe	ed.
Vascline, White 80 g		c. Batch should be allowe	d to cool
Water 60 g	ę.	without stirring to 105° 1	
Transparent, soft, white cream.	- 1	temperature it is poured i	nto jurs.
No. 2	1		
Cetyl Alcohol 10 g	or.	Ob sorin Cold Croum	
Paraffin, Liquid 40 g	or.	Glycerin Cold Cream	
Vaseline, White 50 g		a. Wax, White	80 g.
Water 60 g		Spermaceti Peanut Oil	80 g. 300 g.
No. 3		Vaseline	300 g.
Cetyl Alcohol 10 g	σ.		000 g.
Paraffin, Liquid 40 g	g.	Melt.	
Vaseline, White 15 g	g.	b. Glycerin	120 g.
Water 35 g		Water	120 g.
No. 4	<u> </u>	Borax	10 g.
Cetyl Alcohol 20 g	σ.	Warm up to 90°; pour into	melted a.
Paraffin, Liquid 20 g		Add when cool:	
Vaseline, White 60 g		Perfume Composition, Fre	esh
Water 60 g		Odor	20 g.
In place of the liquid paraffin th	here		•
can be used a good vegetable oil.	The		
maximum water-content (37.5%) can	ı be	Triethanolamine Cold Ci	ream
increased by adding 10% wool fat.		(Water-Soluble, Liquid	1)
Procedure: Melt the fatty mater	rials	g. Paraffin, Liquid	72 g.
together and stir, then run in boil	ling l	Triethanolamine Stearate	14.5 g.
water, a little at a time, not adding	ad-	Dissolve, warming gentl	у.
,,,	_	, 69	-

b. Water, Distilled c. Perfume	160 g. 1.5 g.	
When a is dissolved by wand add b slowly. Let star add perfume, then fill into	arming, stir, id 24 hours,	
Cleansing Cream		
(Semi-Absorbent) [
Lanolin	22 g.	
White Mineral Oil	25 g.	tl
White Petroleum Jelly	11 g.	v
Distilled Water	42 g.	sl
Perfume	to suit	et
Cleansing Cream		ei
(Non-Absorbent)		n w
,		u
Ceresin White Mineral Oil	18 g. 81 g.	c
White Petroleum Jelly	1 g.	
Perfume	0.5 g.	
	_	
Nourishing Creat	m	1
White Beeswax	9 g.	
Spermaceti	3 g.	ĺ
White Petroleum Jelly	35 g.	1
Benzoated Lard	18 g. 4 g.	l
Lanolin Liquid Paraffin	9 g.	
Distilled Water	21 g.	t
Borax	1 g.	e
	-	v
Liquid Nourishing C		a
Lanolin, Anlıydrous	16 g.	1
Stearic Acid	3 g.	1 ^
Tricthanolamine	1 g. 80 g.	1
Water, Distilled	ou g.	
Non-Irritating Cre	e me	1
U. S. Patent 1,979		1
Formula No. 1		1
Vanishing Crea		1
		1
Stearic Acid Lanolin (Anhydrous)	220 g. 40 g.	1
Lanolin (Anhydrous)	40 g.	1

Water 500 g.

The cream is prepared by melting the acid and lanolin and adding them with constant stirring to the remaining ingredients, which are heated to 95° C. An emulsion forms at once which thickens upon cooling. Efficient agitation of the mixture is essential to obtain a smooth product. The solid content, i.e., in No. 1, the lanolin and stearic acid, of a cream of this type may vary from 15% to 35% depending upon the ingredients used and the type of product desired.

Triethanolamine

Ether

Diethylene Glycol Mono-ethyl

12.5 g.

No. 2 Cleansing Cream

Stearic Acid	122.5	g.
Lanolin (Anhydrous)	3 5	g.
White Mineral Oil	210	ġ.
Triethanolamine	17.5	g.
Diethylene Glycol Mono-ethyl		
Ether	40	g.
Water	420	g.

The method of preparing this cream is the same as that employed in the previous formula. A cream of this type should have a fairly high content of the ethanolamine in order to completely emulsify the oil so that it may be removed from the skin by washing with water. Various oils and waxes may be used in this type of cream, and the oil content should be fairly high.

No. 3 After Shaving Cream

Stearic Acid	15	g.
Triethanolamine	0.75	g.
Diethylene Glycol Mono-ethyl		
Ether	8	g.
Menthol Crystals	0.75	ġ.
Ethyl Alcohol (Anhydrous)	0.5	g.
Water	75	g.
The cream is prepared ac	cordin	g t

The cream is prepared according to the procedure given above. In general, creams of this type are similar to the vanushing creams with the addition of an emollient or a medicant, such as menthol, bay rum, witch hazel or the

No. 4

Latherless Shaving Cream

Latherless Shaving Cre		
Stearic Acid	350	g.
Lanolin (Anhydrous)	67.5	g.
White Mineral Oil	169	g.
Triethanolamine	34	g.
Sodium Tetraborate	34	g.
Diethylene Glycol Mono-ethyl	l	
Ether	22.5	
Water	1170	**

This preparation may be made by the procedure given in No. 1 and the oil may be included in the melted acid and wax mixture which is then added to the other ingredients.

Massage Cream

TITTED OF COURT	
White Beeswax	12.5 g.
Paraffin Wax	10 g.
White Mineral Oil	50 g.
Distilled Water	26 g.
Borax	1 g.
Perfume	0.5 g.

Massage Preparations

These substances are dispensed in ointment, mixture or solution form, and ap-

plied before or	treatment,	usually
with a vibrator.		

Formula No. 1	
Menthol	2.5 g.
Tragacanth	4 g.
Glycerin	12 cc.
Alcohol	15 cc.
Water	300 cc.
No. 2	
Gelatin	2 g.
Water	48 cc.
Glycerin	5 cc.
Glycerite of Boroglycerin	45 g.
No. 3	
Fluid Extract of Hamamel	is 10 cc.
Wool Fat	60 g.
Petrolatum	30 g.
No. 4	o
Menthol	0.8 g.
Camphor	0.8 g.
Eucalyptol	3 g.
Petrolatum	96 g.
1 Ctiviatum	. 6

Almond Hand-Cleansing Paste

The "Almond Bran" is made out of two equal parts of sweet and bitter Almonds. One can make a "Glycerin Paste" or a "Camphor Paste."

Glycerin Type

Two hundred fifty pounds of the bran are pounded with 5 lb. of rose water and mixed with the following:

One-quarter pound bean or cornflour, 1-2 chicken eggs, 15 lb. borax, 5 lb. fine potassium carbonate, and about 50 lb. glycerin.

The Camphor Paste is made by adding to the pounded "Almond Bran" a mixture of 25 lb. each of 10% camphor oil and spermaceti, molten together.

After cooling, add a powderized mixture of 100 lb. potato flour and 50 lb. talc, and 100 lb. rose water. Mix well altogether. Color with alkannin or curcuma.

Grind a and b separately, mix, warm then on the water bath until odor of alcohol disappears.

Glycerin-Honey Jelly	,	
Honey	20	g.
Water	500	
Glycerin	450	g.
Agar Agar, Cut	15	g.
Methyl p-Hydroxy benzonte	1	g.
	and	solu-
Warm to complete swelling		
tion percolate, if necessary.	om,	and
add:		
Formaldehyde (40%)	1	g.
Perfume Composition	1	g.
Protective Hand Crean	118	
Formula No. 1		
Zine Stearate, U.S.P.	10	g.
Alummum Subacetate Solu-		
tion N.F. (7½-8%)	15	g.
Gum Camphor Menthol Crystals	3	\mathbf{g}
Menthol Crystals	1,	g.
Acid Carbolic, U.S.P.	1/2	g.
Glycerin, U.S.P.	1/2	g.
Lanolin, Anhydrous	41/	g.
Gnm Tragacanth	41/2 18	g.
Soap (Low Alkalı Content)	10	g.
White Rose Oil Technical	1/2	g.
Triethanolamine	46	g.
Water No. 2	10	g.
*	••	
Zine Stearate, U.S.P.	10	g.
Aluminum Subacetate Solu-		_
tion N.F. (71/2-8%)	15	g.
Gum Camphor	3 1	g.
Menthol Crystals	1/2	g.
Acid Carbolic, U.S.P.	1/2	g.
Glycerin, U.S.P.	1/2	g. g.
Lanolin (Anhydrous)	41/2	
Gum Tragacanth Soap (Low Alkali Content)	18	g.
White Rose Oil Technical	1/2	g.
Triethanolamine	1/2	g.
Water	441/4	g.
Sulpho Ammonium	,-	
Ichthyolato	2	g.
No. 3		
White Rose Technical Oil	35	g
Paraffin Wax	55	g.
Ammonium Sulpho-Ich-	2	~
thyolate	1	g.
Stearic Acid	1/2	g.
Tricthanolamine	71/2	
Water	1 72	ο.
No. 4	_	11
Glyceryl Monostearate		lb.
Magnesium Stearate		lb.
Beeswax		lb.
Petrolatum Munoral Oil White		lb. lb.
i Minoral IIII Walte	υ	

Mineral Oil, White

Water

60 lb.

			L FORMULARY		
	Cuticle Softener	•	Distilled Water	7.875 g.	
	Formula No. 1		Perfume Oil	1.125 g.	
White :	Petrolatum (Short		No. 2		
Fiber		87.75 oz.	Almond Oil	24 g.	
Paraffi	(mp. 125° F.)	9 oz.	Lanolin	22 g.	
Mentho		3 oz. .25 oz.	Soft Paraffin	11 g.	
Thymol		to suit	White Beeswax Rose Water	3 g.	
Color (Oil Soluble Red) No. 2	to suit	Perfume	to suit	
	(Anhydrous)	12 oz.			
	(Distilled)	12 oz. 0.5 oz.	Mosquito Repelling	Cream	
Lecithi	n Petrolutum (Short	0.5 0z.	Formula No. 1		
Fiber		55.5 oz.		5 g.	
	Oil (White)	20 oz.	a. Wheat Starch		
Perfum		to suit	1 (llane in coop DC)	10 g.	
			b. Glycerin (28° Bé.)	45 g.	
	Skin Cream		c. Lanolin d. Clove Oil	30 g.	
a. Stea		85 m	ł	5-10 g.	
Land		85 g. 5 g.	Grind a until homogeneou		
	l Alcohol	10 g.	warm gently until a homo is formed. Cool, and grind	geneous jei Laow with	
	elt together.	6.	and d in a mortar very the		
	erin (28° B6.)	36 g.	distribution is satisfactory.		
	thanolamine	5 ec.	into collapsible tubes.	111 40 011	
Borr		ifepointful	No. 2		
Wat		250 cc.		F 0	
Boil.			a. White Wax	50 g.	
Add b	slowly to a, stir unt	il cold. Per-	Spermaceti	50 g.	
lume as e	lesired is added at	the end.	b. Borax	4 g.	
			7. Ammonia (0.96)	40 g.	
""	enetran'' Skin Co	smetic	(Water	510 cc.	
Paraffi		20 cc.	c. Wheat Starch	1 g.	
	(Whale) Oil	25 сс.	Genera	4 g.	
Parach	ol (Absorption Bas	e) 5 g.	Sodium Benzoate	0.5 g.	
Cholest	erin	0.5 g.	Make up cream as usual r		
Lecithi		2.5 g.	a, then add the solution c v		
Fatty	Oil, Preserved	47 cc.	made up before (soak cold,	then warm	
			clear solution, if necessary, pour through a fine sieve), stir thoroughly, stop hea		
Wri	nkle ''Removing''	Creams	ing, stir until cooled, and a		
Lanolii weight).	cocon butter 10, ste	arin 10. olive	Eucalyptus Oil	50 cc.	
oil 12, ch	olesterol 2, lecithin	4, water 60,	No. 3		
moldex 0	olesterol 2, lecithin 4, sodinm benzoate	e 1. Accord-	Eucalyptus Oil	0.5 cc.	
ing to a	iother method, a m	ielted base is	Caryophyllum Oil	0.5 cc.	
	pared with whi		Lavender Oil	0.5 cc.	
	spermaceti 10, ster		Quinine Sulphate	1 g.	
	coa butter 40, and : In this melt are		Olyceriu Salve to mal	te 100 g.	
	olesterol, with furt		No. 4		
	plete solution, of		Tragacanth To. 4	3 g.	
	dium benzoate and		Alcohol	5 g.	
	ig stirred until it t		Seap Solution	2.5–25 g.	
			Glycerin	45 g.	
	Skin "Food"		To this cream add:		
	Formula No. 1		Menthol	1 g.	
Lanoli	n (Anhydrous)		Sodium Benzoate	ī ģ.	
U.S.	Ρ.	36.4 g.	Citronella Oil	1 cc.	
Sperm	aceti, U.S.P.	6.4 g.	Caryophyllum Oil	0.5 cc.	
Snow	White Petrolatum,		Alcohol Tincture of Green Soap	10 cc. 10 cc.	
U.S.		48.2 g.			

COSMETICS	AN
Mosquito Repellants	
Formula No. 1	1
Pyrethrum Flowers 10 g.	1
Isopropyl Alcohol, or Ethanol	Ī
Isopropyl Alcohol, or Ethanol with Thymol 100 g.	
Oil of Cloves 2 g.	
No. 2	ł
	١.
Oil of Thuia 20 g.	f
Oil of Eucalyptus 45 g. Oil of Thuia 20 g. Oil of Laurel 5 g.	
Phenol 3 g.	
Camphor 20 g.	
Alcohol 100 g.	
Turpentine Oil 50 g.	
Quassia, Tincture 40 g.	
Pyrethrum Extract 50 g	
Xylol to make 1000 cc.	
No. 3	Λ
Pyrethrum Extract 0.5 g.	W
Amyl Saheylate 3.5 g.	
Petroleum (bp. 182-292°;	
sp. gr. 0.801) 96 g.	l
No. 4	
Pyrethrum Powder 1 g.	
Derris-Root Pawder 1 g.	
Tobacco Powder 0.5 g.	
Alcohol, Diluted 25 g.	
Percolate thoroughly and filter; add: oil of eucalyptus or menthol to suit.	
Mosquito Protection Cream	
•	
(Non-Grensy)	١
Formula No. 1 Soak	u
a. Agar-Agar 2 g.	
Water, Cold 400 g.	
Then warm slowly over gentle heat:	
b. Melt Stearin 60 g.	
(Potassum Carbonate 6 g.	
d. Water 410 g.	i
- Glycerin (28° Bé.) 68 g.	
Make up emulsion by warming and	
stirring.	
Add a to the emulsion of b-c in d,	is
Add a to the emulsion of be in d, both should be 80° C; stir continually.	cc
When cold, add 12 g. of the following mixture:	
Cedar Oil 7.5 g.	
Citronella Oil 15 g.	
Camphor 2 g.	
Eucalyptus Oil 4.5 g.	
Alcohol 7 g.	
No. 2	
Treatment as above:	
Agar-Agar 2.2 g.	
Stearin 60 g.	
Potassium Carbonate 4 g.	
Sal Soda 2 g.	ta
•	

	Alcohol	12 g.	
	Beeswax, White	8 g.	
	Lanolin (Anhydraus)	8 g.	
	Glycerin	60 g.	
	Water	830 g.	
	Beta Naphthol	l g	
	Essential Oils as in Forn		
	Treatment as in No. 1, fats (wax, lanolin, stearin)		
ı		together.	
	No. 3		
ı	a. Agar-Agar	2.5 g.	
1	Glycerm	100 g.	
-	Water	750 g.	
	b. Glyceryl Monostearate	120 g.	
ı	Spermaceti Melt.	100 g.	
1	Pour a hot into b, make en	sulaion atie	
1	Add boiling water up to 9		
ļ	when cold:	,	
1	Moldex or Other Good		
1	Preservative	2 g.	
1	Essential Oils	2 g. 12 g.	
1	(See Formula Na,	1)	
1			
1	All Weather Crea	nı	
1	Stenric Acid a. Adeps Lanae, Anhydrou	210 g.	
١	a. Adeps Lanae, Anhydrou	в 50 g.	
1	(Glycerin	133 g.	
١	b. { Glycerin Tricthanolamino Borax	20 g.	
1	O. Borax	5 g.	
١	Distilled Water	582 cc.	
١	Melt up a to about 65° C	, add b boil-	
1	ing bot, in thin jet, stirring	thoroughly	
	until cold.		
١		_	
1	Night Cream (Grea		
1	Parathn Oil, White	2500 g.	
ı	a, Wax, Scale	500 g.	
1	a. Wax, Scale Beeswax, Bleached Adeps Lanae, Anhydrou	500 g. is 500 g.	
1	Adeps range, Annyarou	2000	
1	Distilled Water	3000 cc.	

a. Paratin Oil, White Wax, Scale Beeswax, Bleached Adeps Lanae, Anhydrous	2500	
Wax, Scale	500	g.
a. Beeswax, Bleached	500	
Adeps Lanae, Anhydrous	500	g.
b. { Distilled Water Triethanolamine Borax	3000	cc.
b. { Tricthanolamine	75	g.
(Borax	35	g.
Melt a tagether at 75° C.; a	dd b	which
is at same temperature, to a. cold.	Stir	untıl

Non-Greasy Cream Formula No. 1

	(Stearic Acid	230	g.
a.	Wax, Scale	40	g.
	Stearic Acid Wax, Scale Adeps Lanae, Anhydrous	10	g.
		140	g.
	Glycerin Triethanolamine Borax	13	g.
0	Borax		g.
	Distilled Water	562	cc.

Melt a and warm up b in another conainer. Mix both (a and b should be 65°

C. boiling) pouring b into a in thin jet. Stir until cold.

No. 2

	Stearic Acid	170 g.
a.	Stearic Acid Adeps Lanae, Anhydrous	13 g.
	Wax, Scale	13 g.
	Spermaceti	5 g.
	Wax, Scale Spermaceti Cetyl Alcohol	4 g.
		80 g.
	Glycerin (28° Bé.) Tricthanolamino	13 g.
b. †	Borax	5 g.
	Borax Distilled Water	697 cc.
M	alt up wayes (65-70°)	add b hot

Melt up waxes (65-70°), add b hot (boils) in thin jet, stirring thoroughly. Optionally, 100 water may be substituted by witch hazel (1:1). Stir until cold.

Liquid Cream

	Stearic Acid	50 g.
_	Adeps Lanae, Anhydrous Cetyl Alcohol Beeswax	4 g.
a	Cetyl Alcohol	1 g.
	Beeswax	1 g.
	(Glycerin	20 g.
	Triethanolamine	2 g.
b.	Borax	2 g.
	Witch Hazel (1:1)	75 g.
	Witch Hazel (1:1) Distilled Water	625 ec.

Melt up together a at 60-70° C. Heat b to boiling, then add in thin jet, stirring vigorously, to a. Stir until cold.

To all above-mentioned creams, perfume should be added during cooling (0.5-0.7%). The perfume components should be colorless, and should not irritate the skin. No alcoholic compositions should be used.

Turtle Oil Cream		
1. Diglycol Stearate	14	lb.
2. Mineral Oil	33/4	gal. lb.
3. Lanolin		
4. Petrolatum (White)	2	
5. Water	6	gul.
6. Turtle Oil		fl. oz.
7. Perfume Oil	$5\frac{1}{2}$	fl. oz.
8. Solution Yellow Color		

Made by Dissolving Yellow Dye 2 drams in Mineral Oil 14 fl. oz. 81/4 fl. oz. Method of manufacture:

a. Melt 1, 2, 3, 4, 6 and 8 at 170° F.
b. Heat 5 to 180° F.

c. Add b to a while mixing. Allow mixer to run until batch is completely emulsified. d. Allow batch to cool to 125° F. and

add 7, and mix at low speed.
c. Batch should be allowed to cool without stirring to 100° F. at which temperature it is poured.

Boro-Glycerin Lanolin Cream

Boric Acid	10 g.
a. Glycerin	40 g.
a. Boric Acid Glycerin Water	250 g.
Dissolve.	
b. Lanolin, Anhydrous Vaseline, White	100 g.
Vaseline, White	600 g.
Melt gently.	
c. Rose Oil, Artificial	10 cc.
or Eau de Cologne Oil	20 cc.

Traggeonth-Glycorin Rago (Hgod Rolow)

ragacanti Giyeerii Dase (O.	seu Delow)
Tragacanth, White, Fine	
Powder	1 g.
Glycerin	5 g.
Grind thoroughly in morta	r and add:
Water, Warm	94 g.
Add while stirring and in	small por-
ions, warm up to 40° C.	Stir until

Menthol Cream

paste is homogeneous.

Menthol	0.2	g.
Moldex or Other Good Pre-		_
servative	0.2	g.
Perfume Oil	0.3	
Alcohol (95%)	5	
Dissolve and add		
Glycerin	5	g.
Add above made		
Tragacanth-Glycerin	100	ø.

Lemon Juice Cream U. S. Patent 1,990,676

Five parts of oxy-cholesterin and 95 parts of petrolatum are thoroughly mixed to form an absorption base. Twenty parts of petrolatum and three parts of beeswax are melted together, and 30 parts of the base are added with thorough stirring. Fifty parts of nat-ural lemon juice are added to the above mixture while still hot and stirring is continued until the mass is cool.

Ink Removing Cream U. S. Patent 1,968,304

A substantially non-aqueous cream for the removal of ink stains from the skin contains about 500 g. of zinc stearate, about 300 g. of citric acid, about 500 cc. of 95 per cent ethyl alcohol and about 2000 cc. of diethylene glycol.

Deodorant Cream Formula No. 1

Benzoic Acid	4 g.
Zinc Oxide	12 g.
Lanolin	4 g.
Petrolatum (Snow White)	80 g.
Perfume	to smt
No. 2	

British Patent 425,059

Coconut Oil	63 g.
Lemon Oil	52 g.
Boric Acid, Powdered	21 g.
Starch, Powdered	10.5 g.
Lanolin	0.2 g.
Perfume	0.1 g.
No. 3	
Formaldehyde	1 oz.
Vanishing Cream	99 oz.

Powder Cream Base

Vanishing Cream

Using specified quantities, preparation of the cream base may proceed on the following lines: A mixture of about 500 g. distilled water, 20 g. potassium carbonate and 125 g. glycerin is heated almost to boiling point in a capacious ves-sel constructed of well enamelled mate rial. Two hundred grams steame acid melted in another vessel are cautiously introduced, a little at a time, into the hot potassium carbonate solution. Violent carbon dioxide evolution ensues and continues until the last portion of stearic acid has been added. When gas development ceases, indicating completion of the reaction, heating is discontinued and the batch transferred to another vessel fitted with stirring gear. An additional 1000 g, water and 125 g, glycerin are added and the mix stirred until cold and viscous. Cold-stirring is important for securing a fine, uniform emulsion and for preventing settlement of stearic acid particles. Certain variations in preparaparticles. Certain variations in prepara-tion can be practiced, such as replacement of glycerin by white liquid paraffin or addition of 125 g. groundnut oil to facilitate emulsification.

Ruggles' Cream

Powdered Stearie Acid	75 g.
Potassium Carbonate	15 g.
Distilled Water	320 g.
Powdered Borax	5 g.
Quince Jelly	75 g.
Distilled Water	100 g.
Powdered Zinc Oxide	10 g.
Glycerite Starch	400 g.

Melt the stearic acid. At the same time dissolve the potassium carbonate in 320 cc. of distilled water and heat to

about 170° F. on water bath. Bring stearic acid to the same temperature and mix them. Continue this temperature on the water bath, with occasional stirring, until the reaction is perfectly complete.

Dissolve the powdered borax in 100 ec. of distilled water, add the quince jelly and heat on water bath to about 170° F. Add this mixture to the first, which should be at the same temperature, and again leave on water bath until reaction is complete.

Heat the glycerite of starch to the same temperature, stir in the powdered zinc oxide with a glass stirring rod and add to the other mixture, stirring occasionally.

Let cool and add perfume (oil ylang vlang recommended).

The most unportant essential is to employ a perfect glycerite of starch. Use Kingsford's or other suitable grade of colu starch and U. S. P. Glycerin and make it up fresh for each batch.

It is also essential to have all three batches at exactly the same temperature when mixing them.

Skin Oil with Isocholesterin

Paraffin Oil plus Preserved Fatty Oil	97 ce.
Isocholesteriu, Technically Pure	3 g.
or Same, Chemically Pure	2 g.

Skin Oil with Lanolin Lanoliu, Bleached Paraffin Oil or Fatty Oils 95 cc.

Skin Oil with Wool Wax Wool Wax, Bleached, Purified 5 g. 35 ee Fatty Oil Paraffin Oil 60 cc.

Skin Oil with Cetyl Alcohol Cetyl Alcohol, Fure Paraffin Oil plas Fatty Oil, 97-95 cc. Cetyl Alcohol, Pure Preserved (1:1)

Skin Oil with Triethanolamine Oleate Triethanolamine Oleate, Pure 2 g. 98 cc. Fatty Oil

Non-Irritating Skin Oil Diglycol Laurate Neutral 4 g. 96 ec. Olive Oil Perfume to suit

Lecithin Skin Oil	Witch Hazel Skin Oil		
Formula No. 1	Witch Hazel Leaves, Powder 100 g.		
Lecithin from Eggs 10-30 g.	Fatty Oil, Preserved 900 cc.		
Paraffin Oil 170-190 cc. Olive Oil. Preserved 800 cc.	Pour hot oil over leaves, let stand for 8 days. Filter.		
Olive Oil, Preserved 800 cc. Perfume, to suit 5 g.	8 days. Filter.		
No. 2	Massage Oil		
	Paraffin Oil 75 cc.		
Lecithin from Brain Sub- stance 20 g.	Parachol (Absorption Base) 5 g.		
Paraffin Oil 180 cc.	Parachol (Absorption Base) 5 g. Olivo Oil, Preserved 20 cc.		
Olive or Peanut Oil, Pre-			
served 800 cc.	Muscle Oil		
Skin Oil "Huile Ambrosiaque"	Castor Oil, Deodorized 66.6 cc.		
Ambergris, Best Quality 10 g.	Alcohol (92-95%) 33.3 cc. Cholesterin, Pure 0.1 g.		
Behen Oil 990 cc.			
Perfume to suit	Sport Oil (for Swimmers)		
Grind the amber with glass powder	Octadecyl Alcohol (Pure) 5 g.		
and introduce into the warmed oil. Shake well. Filter after 3-4 weeks.	Fatty Oil, Preserved 55 cc.		
Well. Illied dittel o I weeks.	Paraffin Oil 40 cc.		
CIL CIL III I I I I I I I I I			
Skin Oil with Wool Fat Alcohols	Cholesterin Oil		
Parachol (Absorption Base) 5-10 g.	Fatty Oil, Pure, or in		
Paraffin Oil 95-90 cc.	Mixture with Paraffin Oil 1000 cc. Cholesterin, C.P. 5-10 g.		
Skin Cleansing Oil	Cholesterin-Lecithin Oil		
Parachol or Absorption Base 2 g.	Same as Cholesterin Oil, but besides		
Triethanolamine Oleate, Pure 0.5 g.	add Lecithin (Eggs, Brain-Substance)		
Fatty Oil, Preserved 97.5 ec.			
	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine.	add Lecithin (Eggs, Brain-Substance) 20-30 g. Face Lotions		
Fatty Oil, Preserved 97.5 cc. Add a little Tricthanolamine. Skin Nourishing Oil	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Trietlanolamine. Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g.	add Lecithin (Eggs, Brain-Substance) 20-30 g. Face Lotions Formula No. 1 Triethanolamino 0.5 cc.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g Paruchol (Absorption Buse) 5 g Lecithin 1 g.	add Lecithin (Eggs, Brain-Substance) 20-30 g. Face Lotions Formula No. 1 Triethanolamino 0.5 cc. Glycerin 4 cc.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genu-	add Lecithin (Eggs, Bran-Substance) 20-30 g. Face Lotions Formula No. 1 Triethanolamino 0.5 cc. Glycerin 4 cc. Alcohol 33 cc. Distilled Water 62 cc.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genu-	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Paruchol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Paruchol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil 5 g. Paruchol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Pedodorized 20 cc. Fatty Oil, Preserved 69 cc.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Peodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g.	add Lecithin (Eggs, Brain-Substance) 20-30 g. Face Lotions Formula No. 1 Triethanolamino 0.5 cc. Glycerin 4 cc. Alcohol 33 cc. Distribed Water 62 cc. Perfumo 0.5-1 cc. No. 2 Triethanolamine 0.5 cc. Glycerin 4 cc. Alcohol (30%) 95.5 cc. Perfume to suit to suit		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Paruchol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesterin, Artificial 3 g.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesterin, Artificial Fatty Oil (Oilve, Sesame,	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesteria, Artificial Fatty Oil (Olive, Sesame, Peanut), Preserved 92 cc.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil Parachol (Absorption Base) Sperm (Whale) Oil, Genuine, Deodorized Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) Oxycholesterin, Artificial Fatty Oil (Olive, Seaame, Peanut), Preserved 92 cc.	add Lecithin (Eggs, Bran-Substance) 20-30 g. Face Lotions Formula No. 1		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Paruchol (Absorption Buse) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesterin, Artificial 7 g. Fatty Oil (Olive, Sesame, Peanut), Preserved 92 cc. No. 2 Parachol (Absorption Base) 5 g.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil Parachol (Absorption Base) Sperm (Whale) Oil, Genuine, Deodorized Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) Oxycholesterin, Artificial Fatty Oil (Olive, Seaame, Peanut), Preserved 92 cc.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesterin, Artificial 3 g. Fatty Oil (Olive, Sesame, Peanut), Preserved 92 cc. No. 2 Parachol (Absorption Base) 5 g. Cetyl Alcohol, Pure 3 g.	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Fatty Oil, Preserved 97.5 cc. Add a little Triethanolamine. Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesterin, Artificial 3 g. Fatty Oil (Olive, Sesame, Peanut), Preserved 92 cc. No. 2 Parachol (Absorption Base) 5 g. Cetyl Alcohol, Pure 3 g.	Add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesteria, Artificial Fatty Oil (Olive, Sesame, Peanut), Preserved 92 cc. No. 2 Parachol (Absorption Base) 5 g. Cetyl Alcohol, Pure 3 g. Fatty Oil, Preserved 91 cc. Astringent Skin Oil	add Lecithin (Eggs, Brain-Substance) 20-30 g.		
Skin Nourishing Oil Egg Oil 5 g. Parachol (Absorption Base) 5 g. Lecithin 1 g. Sperm (Whale) Oil, Genuine, Deodorized 20 cc. Fatty Oil, Preserved 69 cc. Skin "Stimulating" Oils Formula No. 1 Parachol (Absorption Base) 5 g. Oxycholesterin, Artificial Fatty Oil, Oilow, Sesame, Peanut), Preserved 92 cc. No. 2 Parachol (Absorption Base) 5 g. Cctyl Alcohol, Pure 3 g. Fatty Oil, Preserved 91 cc. Astringent Skin Oil	Add Lecithin (Eggs, Brain-Substance) 20-30 g.		

No. 5		Alcohol	6 g.
Triethanolamine	5 cc.	Glycerin, C.P.	3 g.
Alcohol (96%)	500 cc.	Almond Oil	10 g.
Spirits of Camphor	100 cc.	Distilled Water ab	out 85 g.
Perfume	10 cc.		-
Glycerin	20 cc.	Face Lotion (For Oi	ly Skin)
Witch Hazel, Distilled		Sulphur, Precipitated	2 g.
		Glycerin, C.P.	5 g.
For Dry Skin: No		Camphor Spirit (10%)	3 g.
Mineral Oil, White	35 cc.	Lavender Water	10 g.
Beeswax	20 g.	Borax	1 g.
Amino Stearin	8 g.	Distilled Water	81 g.
Water	50 cc.	Distilled water	0.1 B.
Warm together and mix	vigorously		•
ntil emulsified.		Acne Face Lot	ion
No. 7	l	Formula No.	1
	72 cc.		_
Vaseline Oil	14 g.	Acetic Acid (96%) or	κ
Amino Stearin	200 cc.	Benzoie Acid	5 g. 500 g.
Water	200 cc.	Alcohol (95%)	300 g.
No. 8		Lavender Oil	4 g.
Triethanolamine	5 cc.	Water	466 g.
Aromatic Spirit	30 ec.	Glycerin (28° Bé.)	25 g.
Bergamot Öil	12.5 ec.	Let stand several weeks.	Filter.
Oil Orange Flowers	0.5 cc.	No. 2	
Lemon Oil	2 cc.		
Rosemary Oil	15 cc.	Potassium Soap from	100 ~
Alcohol (70%)	940 cc.	Olive Oil (Neutralized)	100 g.
No. 9	ľ	Alcohol (90%)	500 g.
	25 g.	Lavender Oil	5 g.
Camphor	850 ce.	Rose Oıl, Artificial	5 g.
Alcohol	25 cc.	Water	390 g.
Glycerin	30 cc.		-
Perfumo Mixture	1570 cc.	Face Water	
Distilled Water	10,0 00	Tricthanolamine	0.5 g.
No. 10	. 1	Glycerin	4 g.
Boric Acid	10 g.	Alcohol	33 g.
Glycerin	29 cc.	Perfume	0.5 g.
Menthol	1 g.	Distilled Water	62 g.
Perfume	5 cc.	Distinct water	В
Alcohol	255 cc.		-
Hamamelis Distillate	300 cc.	Prophylactic Face	Waters
Rose Water	400 cc.	Formula No.	1
No. 11		Ammonium Chloride, C.	
Alcohol	450 cc.		20 cc.
Camphor, Spirits of	100 cc.	Witch Hazel	10 cc.
Perfume	10 cc.	Rose Water	69.5 cc.
Hamamelis Distillate	440 cc.	Distilled Water	00.0 66
		No. 2	
No. 12	400	Ammonium Chloride	2.5 g.
Potassium Carbonate	400 g.		10 ec.
Distilled Water	2000 cc.	Cherry Laurel Water Witch Hazel	10 cc.
Orange Flower Water	1000 cc.	Rose Water	20 cc.
Alcohol	100 cc.	Distilled Water	57 cc.
Perfume	to suit	Diethylene Glycol	0.5 cc.
No. 13		Dietayiene dijeoi	
Borax	50 g.		-
Sodium Thiosulphate	500 g.	Kummerfeld's (Fac	e) Water
Distilled Water	8500 cc.	Sulphur, Colloidal, or I	Pinely
	500 сс.	Descipitated	2 g.
	500 cc.	Precipitated	12 cc.
Glycerin		Glycerin	4 cc.
Eau de Cologne			
Eau de Cologne		Spirits of Camphor	
Eau de Cologne	y Skin)	Eau de Cologne	20 cc.
	y Skin) 0.05 g.	Spirits of Camphor Eau de Cologne Distilled Water Optionally: Addition of	20 cc. 100 cc.

sh, or Triethanolamine (iect).	intensifies ef-	Skin Hardener Alum	30 g.
		Water and Alcohol (Equal	0
Sulphur Face Wa	ater	Parts)	250 cc.
Sulphur, Colloidal	3 g.		
Potassium Carbonate	1.5 %.	Strong Astringent Lo	tion
Glycerin	1.5 g. 5 ec.		
Spirits of Camphor	4 cc.	Salicylic Acid	3¼ lb.
Alcohol	10 сс.	Benzyl Cinnamate	21/2 oz.
Distilled Water	76.5 cc.	Acetone	1 gal.
Distinct Water	1000 001	Alcohol	1 gal.
		The quantity of salicylic a	icid may h
Skin Lotion		reduced 1/2 if a milder agent	
Gum Tragacanth	4 oz.		
Glycerin	3 oz.	Face Water with Witch	Hand
Phonol	1 oz.		
Oil of Teel Water	120 oz.	Alcohol (40%)	920 g.
Water	360 oz.	Witch Hazel	50 cc.
Perfume	2 oz.	Alcohol (40%) Witch Hazel Glycerin, C.P.	30 g.
Modern Glycerin-Sulphu	r Lotion	Modern Neutral Face	Water
		Alcohol (40%)	920 cc.
Colloidal Sulphur in Glyc	erin 100 -	Diethylene (Hycol	30 g.
(24%)	100 g.	Diethylene Glycol Glycerin, C.P.	50 g.
Tineture of Green Soap	100 g.		0° 6.
Eau de Cologne—Oil Water, Distilled	1 g.	Dans Water for Mottled Chin	an Massiele
Water, Distilled	799 g.	Face Water for Mottled Skin	
		Zinc Sulphate, C.P.	1 g. 0.5 g.
Glycerin and Cucumber	r Lotion		0.5 g.
Cucumber Perfume	5 g.	Hydrogen Peroxido	
		(3-10%)	89.5 cc.
b. {Alcohol (95%) Benzoic Acid Cucumber Perfume	ου g.		
b. Benzoic Acid	0.3 g.	Freckle Lotion	
(Cucumber Pertume	υg.	Dissolve:	
o. Tragacanth, Fine, Whi	te 5 g.	Potassium Carbonate	60 g.
Glycerin	100 g.	Potassium Chlorate	20 g.
Grind c together, then ad	da and b in	Borax	15 g.
mall portions, gunding to a	get homogene-	Sugar	60 g.
us priste.	, ,	In:	J
•		Rose Water	220 ~
Consulting and Non-1			330 g. 355 cc.
Cucumber and Egg 1		Orange Flower Water Glycerin	150 cc.
Cucumber Juice	400 g.	Giyeeiin	100 сс.
Alcohol	50 g.		
Alcohol Benzoic Acid Egg Yellow Lavender Oil Rose Oil, Artificial Glycerin	0.25 g.	Skin Cleansing Loti	on
Egg Yellow	1-2 g.	British Patent 423,4	26
Lavender Oil	3 g.	G. Hom Dilomete	1 22
Rose OII, Artificial	1 g.	Sodium Biborate	1.33 g.
Glycerin	100 g.	Potassium Alum	2.30 g.
		Soda Ash Water	1.75 g. 100 cc.
Face Water, Ac	id	1	
		Evaporate down to half of	volume.
Alcohol (45%) Tri- (or Di-) Ethylene Gl	veol 30 cr		
Citric Acid	5 g.	Liquid Deep Pore Cle	anser
Glycerin	30 g.	Witch Hazel Extract, U.S.I	P. 50 oz.
Witch Hazel	35 cc.	Alcohol	28 oz.
TITOM AIRECT	00 00.	Polyalkyl-glycol Ether	
		(Glycopon S)	15 oz.
Face Water, Astric	igent	(======================================	
Alcohol (35%)	950 cc.	Face Pack	
Diethylene Glycol	30 g.		
a	15 g.	Put on face for 20 min. a	mixture o
Glycerin			
Alcohol (35%) Diethylene Glycol Glycerin Tannic Acid, Pure Phosphoric Acid, C.P.	3 g. 2 g.	Oat Flour	20 g.

The same same same same same same same sam		
Chamomile Flowers		2 o.
Hamamelis Leaves		2 g. 2 g. 2 g.
Rosemary Leaves		2 g.
Camphor Water) ec.
Treat afterwards with as		
tion of	ti ingt	110
Tannic Acid	0.2	5 g.
Rose Water	25	g.
Hamamelis Water	50	g.
Orange Flower Water	25	g.
Hand Lotion		
Formula No. 1		
Alcohol, Ethyl		ec.
Glycerol	100) ee.
Menthol	:	σg.
Perfume, Rose Oil, Etc.		cc.
Salicylic Acid		2 g.
Water	300	ee.
No. 2		
Alcohol, Ethyl		ec.
Glycerol		ce.
Menthol	:	} g.
Perfume, as desired, about	1	cc.
Salicylic Acid	2	2 g.
Water	273	ec.
No. 3		
Alcohol, Ethyl	500	ec.
Glycerol	250	ce.
Menthol	1	g.
Perfume, as desired, about	1	ee.
Salicylie Acid	:	2 g
Water	250	ie.
A lavender coloration of	varvi	ng in-
tensity may be obtained by ac	ding	traces
of ferric chloride solution. F	ormu	la No.
OF THE CHIOTEST INSTITUTE I		

Low Cost Almond Lotion 1. Diglycol Stearate 7 lb. gal. 30 2. Water

3. Gum Tragacanth Sogal, fl. oz. lution 3 4. Benzaldchyde 11/2 fl. oz. 5. Oil of Bergamot

Method of manufacture:

3 gives a rather only lotion.

a. Melt No. 1 at 160° F. b. Heat No. 2 to 205° F. and run into stone jar (note final temperature of water after dumping into jar must not be below 170° F.).

c. With high speed agitator running, add a (molten at 160° F.) to b, at at least 170° F. and allow mixer to run until temperature has dropped to 140° F.

d. Add 3 to batch while mixture is still running.

c. Add 4 and 5 immediately after 3 and allow mixer to continue running until temperature has dropped to 90° or 95° F.

The gum solution is made as follows: Gum Tragacanth 21/2 lb. 50 gal. Water

Allow the gum to soak for several hours and beat into solution.

Rose Lotion

1. Diglycol Stenrate 114 2. Water 30 gal. 3. Gum Solution 6 gal. fl. oz. 4. Oil of Rose Solution 5. Red Color

Made by Dissolving Red Dye, 1 oz., in

3/4 fl. oz. Water, 1 qt.

Method of manufacture: a. Melt No. 1 at 160° F.

b. Heat No. 2 to 200° F. and run into stone jar (note: final temperature of water after damping into jar must not be below 170° F.).

c. With high speed agitator running add a (molten at 160° F.) to b at at least 170° F. and allow mixer to run until temperature has dropped to 140° F.

d. Add 3 to batch while mixer is still

running. c. Add 4 and 5 immediately after 3 and allow mixer to continue running until temperature has dropped to 90° or 95° F.

The gum solution is made as explained under almond lotion.

Lemon Lotion

1. Diglycol Stearate 2. Water 30 gal. 3 Gum Solution gal. 4. Oil of Lemon 5. Yellow Dye 3/4 oz.

Method of manufacture:

a. Melt No. 1 nt 160° F. b. Heat No. 2 to 200° F. and run into stone jar (note: final temperature of water after dumping into jar must

not be below 170° F.).

c. With high speed agitator running add a (molten at 160° F.) to b at at least 180° F. and allow mixer to run until temperature has dropped to 145° F.

d. Add 3 to batch while mixer is still running.

e. Add 4 and 5 immediately after 3 and allow mixer to continue running until temperature has dropped to 95° or 100° F.

The gum solution is made as explained under almond lotion.

76 THE	CHEMICA	L FORMULARY		
Milky Lotion with Pe Base Emulsion (See Below		Borax c. Borax	20	g.
Distilled Water Perfume Base Emulsion	445 g. 5 g.	Water, Warm Add c cold to a and b.	880	g.
Distilled Water	710 g.	Dusty Odor Face Lo	tions	
Mineral Oil Dried Pectin	180 g. 50 g.	Formula No. 1		
Citric Acid	10 g.	Glycerin	1	cc.
Extract Chamomile Flowers	50 g.	Lactic Acid Menthol	0.2	cc.
Moisten the pectin with a l		Opoponax—Perfumes with		
and then rub with a little wa		Violet Root Oil, etc.		cc.
		Alcohol (35%)	0.3 97.5	
mucilage is obtained. The p to a large extent. In the	rest of the	No. 2		
water dissolve the liquid charact and the warm solution		Glycerin	1	cc.
time to the pectin mucilage.		Citric Acid Aluminum Acetato	$0.2 \\ 0.3$	g.
the water has been added, l	eat until a	Menthol	0.5	g.
uniform solution results, ave	diffing over-	Hamamelis Water	5	cc.
this solution, preferably in a		Perfumes (as above) Alcohol (40%)	0.5 92.5	
or a homogenizer.		No. 3	02.0	
		Glycerin	1	cc.
Bathing Milk		Alum	1	g.
Emulsion of:		Zinc Sulphophenylate Perfumes (as above)	0.5 0.5	g.
Turkey Red Oil Neutral- ized with Caustic Potash	200 g.	Menthol	0.5	g.
Perfume Mixture	350 g.	1sopropyl Alcohol	10	cc.
Add then:	· ·	Rose Water Alcohol (30%)	10 76.5	cc.
Potassium Carbonate So-				
lution (20° Bé.) Clear Liquid Soap (10%)	50 g. 400 g.	Eau de Quinine		
A higher content of etheric		Alcohol	600	g.
tates more turkey red oil		Water Quinine Sulphate	400 5	g. g.
and eventually terpineol.	1 100 -	Saponine	1	g.
For a thicker balm: Use Turkey Red, but add 100-1	50 g. oleic	Saffron Tincture	2	g.
acid, and saponify the whole		Orseille (Red Dye) Rose Oil	0.2 2	g.
tic. The milky character is bett	ared by ad-	Musk, Tincture	1	g.
dition of potassium stearate		Lemon Oil	1	g.
amine stearate (or oleate).		Eau de Cologno (509	%)	
		Bergamot Oil	10	cc.
Benzoin Milk		Lemon Oil	14	cc.
Mix in a mortar or dish:		Citral Thyme Oil, White	1.4 2.6	
Tincture of Benzoin	50 cc.	Rosemary Oil	3.4	
a. Alcohol (95%)	200 сс.	Lavender Oil	10	
 b. Glycerin c. Water, Distilled 	700 cc.	Ixolene, Extra Alcohol	3.4 500	cc.
First grind a, add b, and	pour slowly	Water		cc.
under stirring c into a and b	Let stand	Cl. The Land		
a week. Filter. Shake befor	e use.	Chypre Head Lotio	n 1.4	00
Glycerin Toilette Wa	tor	Geraniol, C.P. Cedar Wood Oil, Rectified	0.25	
(Alcohol (95%)	50 g.	Benzyl Acetate, Chlorine-		
a. Rose Essence	0.4 g.	Free Hydroxycitronellal, C.P.	0.6	cc.
b. Glycerin	50 g.	(100%)	0.7	cc.

0.25 cc.

Storax Ull	0.25 cc.
Geranium Oil, Réunion	0.6 cc.
Benzyl Benzoate Linalyl Acetate	2.5 cc.
Limple Assets	
Linklyi Acetate	0.8 cc.
Linalool, Extra	1.2 cc.
Anise Aldehyde	0.1 cc.
Iris Oil, Genuine, Concrete	
This Oil, Genuine, Concrete	
Coumarin	0.15 g.
Civet, Genuine (100%)	0.02 g.
Civet, Genuine (100%) Patchouli Oil, Genuine	0.2 cc.
Musk, Artificial,	
"Ambrette"	0.2 g.
	0.2 g.
Musk, Artificial,	
"Ketone"	0.05 g.
Labdanum Extract	0.15 cc.
Vanillin	0.13 g.
Phenylethyl Alcohol	0.6 cc.
Rosemary Oil	0.05 cc.
Alcohol 6	70 cc.
Distilled Water 3	20 cc.
Distinct water	
Alcoholic Sulphur Hair I Sulphur Glycerin Solution (24%) Water	otion 5 g. 20 ec.
Salicylic Acid	0.5 g.
Menthol	0.3 g.
Alcohol (24%)	70 ec.
	to suit
Perfume	
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour	.ssage 62 25 g. 30 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonato Buckwheat Flour Henna	25 g. 30 g. 10 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonato Buckwheat Flour Henna Salicylic Acid	25 g. 30 g. 10 g. 5 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur	25 g. 30 g. 10 g. 5 g. 5 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonato Buckwheat Flour Henna Salicylic Acid	25 g. 30 g. 10 g. 5 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonato Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil	.ssage 62 25 g. 30 g. 10 g. 5 g. 5 g. 5 cc.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Decodorized Kerosene	.ssage 62 25 g. 30 g. 10 g. 5 g. 5 g. 5 cc.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 3 oz.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin	25 g. 30 g. 10 g. 5 g. 5 cc.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 3 oz.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate	25 g. 30 g. 10 g. 5 g. 5 g. 5 cc.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 7 oz.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 7 oz. to 3 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid	80 oz. 3 gc. 10 oz. 10
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid	80 oz. 3 gc. 10 oz. 10
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth.	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 7 oz. to 3 g. Stir in
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid	80 oz. 3 gc. 10 oz. 10
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth.	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 7 oz. to 3 g. Stir in
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth. Alcohol, Anhydrous Hair Setting Concentr	25 g. 30 g. 10 g. 5 g. 5 g. 5 g. 5 cc. 80 oz. 3 oz. 10 oz. 7 oz. to 3 g. 6 g. Stir in 48 g. ato
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth. Alcohol, Anhydrous Hair Setting Concentr Karaya Gum	25 g. 30 g. 10 g. 5 g. 5 g. 5 g. 5 cc. 80 oz. 3 oz. 7 oz. to 3 g.) 6 g. Stir in 48 g. ate 12 g.
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth. Alcohol, Anhydrous Hair Setting Concentr Karaya Gum	25 g. 30 g. 10 g. 5 g. 5 g. 5 g. 5 cc. 80 oz. 3 oz. 10 oz. 7 oz. to 3 g. 6 g. Stir in 48 g. ato
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth. Alcohol, Anhydrous Hair Setting Concentr	25 g. 30 g. 10 g. 5 g. 5 g. 5 g. 5 cc. 80 oz. 3 oz. 7 oz. to 3 g.) 6 g. Stir in 48 g. ate 12 g.
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Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth. Alcohol, Anhydrous Hair Setting Concentr Karaya Gum Glycerin or Glycol Alcohol Perfume	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 10 oz. 7 oz. to 3 g.) 6 g. Stir in 48 g. ate 12 g. 12 g. 12 g. 12 g. to suit
Preparation for Head Ma German Patent 616,3 Lauryl Sulphonate Buckwheat Flour Henna Salicylic Acid Sulphur Castor Oil Scalp Stimulant Deodorized Kerosene Resorcinol Monoacetate Lanolin Diglycol Laurate Hair Wave Concentra Karaya Gum Glycol Bori-Borate (Liquid Rub together until smooth. Alcohol, Anhydrous Hair Setting Concentr Karaya Gum Glycerin or Glycol Alcohol	25 g. 30 g. 10 g. 5 g. 5 cc. 80 oz. 3 oz. 10 oz. 7 oz. to 3 g.) 6 g. Stir in 48 g. ate 12 g. 12 g. 12 g. 12 g. to suit
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Storax Oil

Liquid Hair	Fixative
Tragacanth, Powder	0.2-0.5 g, 5-10 g,
Glycerin, C.P. Alcohol (95%)	1 g.
Distilled Water	93.8-88.5 cc.

Dissolve gum in hot water, adding it together with the glycerin (ground together previously), filter; perfume with water soluble essential oils, or use orange flower (rose flower) water instead of distilled water, then dve pale green.

of tilled water, then dye pale green.

If paste is wanted for collapsible tubes, use 3-4 g. of gum tragucanth.

Brilliantine		
Oil of Bitter Almond	1.5	cc.
Oil of Clove	3	ee.
Oil of Bergamot	6	cc.
Custor Oil	50	ee.
Glyceryl Monoricinolente	50	g.
Suet	50	g.

Non Greasy Brilliantine

Diglycol Laurate	40 cc.
Alcohol	60 ec.
Perfume and Color	to suit

Hair Fixative Creams

The simplest type of fixative cream is a tragacanth nuclage containing up to 25% of liquid parallin, more or less enulsified. Such creams require vigorous shaking, as the oil separates on standing. Permanent creams which now enjoy tremendous popularity, thanks to good advertising and their own inherent good qualities, are of two types:—oil-in-water emulsions and water-in-oil emulsions, the oil in both cases being mainly liquid parallin. The most popular of these new fixatives is of the second type, a water-in-oil emulsion. It is not, as it is often supposed, a tricthanolamine emulsion, but resembles a semi-liquid cold cream. A formula for this type of cream, which has been published and widely quoted, is as follows:

Formula No. 1 Liquid Paraffin

White Beeswax	100 g.
Borax	6 g.
Water	150 cc.
No. 2	
Liquid Paraffin	45 cc.
Stearic Acid	5 g.
Water	49 cc.
Triethanolamine	1 cc.
Parforma	to suit

3000 cc.

Add the liquid paraffin and stearin heated to about 65° C, to the solution of triethanolamine in water at the same

temperature, and stir until it thickens. When nearly cold add the perfume. Avoid too vigorous stirring which causes frothing.

This formula gives a very thick cream which can easily be thinned by diluting with water if desired.

Hair Fixative Perfumes

The popular ingredients include the citrus oils (orange, lemon, bergamot and lime), lavender, rosemary, geranium, petitgrain and coumarin; about 1% of perfume is sufficient. The following table will serve as a guide:

Formula	No. 1	No. 2	No. 3	No. 4	No. 5	No. 6
Bergamot Oil	55 cc.	20 cc.	45 ec.	40 сс.	50 cc.	40 cc.
Lavender Oil	10 cc.	50 cc.		50 cc.	_	40 cc.
Lemon Oil	3 cc.	_	20 cc.			
Orange Oil	5 cc.		5 cc.		15 cc.	
Lime Oil	5 cc.	-	5 cc.		_	
Petitgrain Oil	15 cc.	15 cc.	25 cc.		10 cc.	
Rosemary Oil	5 cc.	5 cc.			5 cc.	
Geranium Oil	2 cc.		_	_	15 cc.	20 cc.
Coumarin		10 g.		10 g.	5 g.	-

Hair Oil	
Formula No. 1	
Alcohol, Ethyl	400 cc.
Glycerol	200 cc.
Perfume, as desired, about	1 cc.
Salicylic Acid	2 g.
Water	400 cc.
No. 2	
Alcohol, Ethyl	400 cc.
Glycerol	300 cc.
Perfume, as desired, about	1 cc.
Salicylic Acid	2 g.
Water	300 cc.
Lavender coloration may	he effecte
by the addition of traces of	ferric chl
ride. The preparation is	completel

nwater soluble, hence readily removed by washing, yet it serves as an excellent "stay comb."

Soapless Shampoo		
(Wetting Out		
)	450 g.	
Oil Oil	50 g.	
	300 0	

Mineral Alcohol to make 1 l. Water Soapless Shampoo Powder

Lohrinol

Agent

Borax Sodium Bicarbonate	oz. oz.
Soda Ash Saponin	0 z. 0 z.
"Oil-Hair Wash"	

Formula No. 1 Diethylaminoethyloleyl Citrate Chamomile Extract cc. Lemon Juice

Alcohol (50%) No. 2 Rape Seed Oil 50 cc. 30 cc. Hazelnut Oil 5 cc. Spike Lavender Oil

Water, Distilled, or

Egg Shampoo

81.5 cc.

Prepare just before use. Separate the yolks and whites of four or more eggs in separate bowls. To the yolks add a tablespoonful of cold water and beat until uniform with an eggbeater. Wash off the beater and beat the whites until fluffy and firm. Add the beaten yolks to the whites and fold the former into the latter. The hair is washed and rinsed with lukewarm water. Then work the egg shampoo, a little at a time, into the scalp and hair. Finally wash and rinse the hair with a strong spray of tepid (not hot) water.

Shampoo Powder

Sulphonated Lorol or 40 g. Lohrinol 40 Sodium Sesquicarbonate

This gives an excellent lather.

Many such additions will suggest themselves to those who wish to experiment. Some people include a specially prepared saponin, 2 to 5%, to help the lather-producing properties.

Liquid Hair Shampoo

Potash Soft Soap Potassium Carbonate

Glycerin	7 g.
Benzaldehyde	0.25 g.
Distilled Water	938 cc.

The procedure is to dissolve the soft soap, with gentle heating, in half the water. The potash, glycerin and benzaldehyde are incorporated in the rest of the water. After the two solutions have been well mixed by stirring, the finished product is left for a week before decanting, filtering and bottling. At first the perfume will be found to disappear, owing to the splitting up of the benzaldehyde into sodium benzonte and benzyl alcohol—but after the lapse of some days the characteristic almond odor will reappear, owing to the oxidation of the alcohol back to the allehyde.

In the above formula, the soap content may naturally be increased if desired—also a proportion of alcohol may be added. Instead of the ahmoud per fume imparted above, a stable fougere or similar compound can be employed. Likewise pine tar, or a 10% solution of henna, may be incorporated in the case of antiseptic or liquid henna shampoos respectively. Novel ingredients for imparting a pleasantly "medicated" odor include iso-thymol.

In the manufacture of liquid soap shampoos, careful control at all points is essential. Turbidity must at all costs be avoided, and for this reason distilled water only should be used and the soap itself completely saponified. Unless proper facilities are available for saponification on the premises, it is better to purchase a ready-made soft soap base (carefully standardized examples of which are now on the market).

Shampoos should, in certain cases, be aged for even longer than a week (e.g., 15 to 30 days), then decanted into a tank fitted with a refrigerating coil, chilled to a low temperature and finally filtered through asbestos. It has been suggested that the period of aging can be radically reduced by first running the shampoo through a colloid mill or homogenizer.

Hair Wash

Liquid Soap Triethanolamine Laurate Alcohol	90-95 10-5 10-5	oz.
---	-----------------------	-----

Hair Washing Soaps Formula No. 1 (for Oily Scalp)

Castor Oil	4,750	
Caustic Potash (50%)	about 7,515	g

Distilled or			
Softened Water	7	6,000	cc.
Perfume, or			
Chanionnle Extrac	t, or		
Wood Tar, Pure, o	r Bette	r	
Perfume Blended			it
Extract	with 500-	2,000	ŧс.

No. 2	·
Coconut Oil	11,000 g
Olive Oil	4,750 g
Canata Datach	,

Caustic Potash		
(50%)	about 7,520 g.	
Distilled or Softened		

Water		76,000 cc.	
Perfume or	Extract	500- 2,000 сс.	
No. 3	(for Dr	y Scalp)	

Coconut Oil	15,000 g.
Olive Oil	6,000 g.
Caustic Potash (50%)	10,200 g.
Glycerin	10,000 g.
Alcohol (95%)	6,000 cc.
Distilled or	

Softened Water 53,000 cc. Perfume or Extract 500- 2,000 cc.

Dandruff Remover

Mercury Bichloride	0.5 g.
Resorcinol	5 g.
Alcohol	125 cc.
Water	125 ec.
Dissolve the bichloride a	nd the resor-
emol in the water. Then	add alcohol.
Apply on the dry scalp a	nd rub thor-
oughly-then shampoo the	hair. One
treatment a week is usually	sufficient for

Dandruff Lotion

a complete absence of dandruff.

Salicylic Acid	2	oz.
Sulphur (Precipitated)	4	OZ.
Castor Oil	10	υz.
Gum Tragacanth	1	oz.
Glycerin "	1	07.
Perfume	0.5	υZ.
Water	82	oz.

Henna, White

Henna white is a bleach, varying in composition with various users. One formula, sodium perborate, 18 g.; henna leaves, 2 g.; affords an excuse for the name. No other excuse can be seen for the waste of henna leaves. Some use

e waste of nenna leaves.	Dune and
Magnesium Carbonate	68 g.
Sodium Perborate	32 g.

Make into a paste a 50-50 mixture of hydrogen peroxide and water before use.

Birch Water

Birch Bud Oil Glycerin	10 40	g.

Soap Spirit	250	g.	Orange Flower Water,		
Ethanol or	250	_	Triple	100 800	cc.
	650	g.	Alcohol	800	cc.
Bergamot Oil	5	g.			
Geranium Oil	1	g.	Eau de Lavende, Am	brée	
Orange Flower Oil	0.5		Lavender Oil, French	50	cc.
Water	50	g.	Lavender Oil, French Bergamot Oil Musk Infusion Ambreine	12	
			Musk Infusion	12	
Florida Water			Ambreine		
	5	cc.	Lemon Oil	6	cc.
Neroli Oil, "Bigarade" Lavender Oil, English Bergamot Oil	5	cc.	Benzoin Infusion	6	cc.
Boronmot Oil	30	cc.	Idola	2	cc.
Limette Oil	2	cc.	Alcohol (96%)	2500	cc.
Clove Oil		cc.	Lemon Oil Benzoin Infusion Idola Alcohol (96%) Water, Distilled	500	cc.
Cassia Oil		cc.			
Cinnamon Oil		cc.	Eau de Cologne		
Rose Oil	5	cc.	,		
Ambra, Liquid, Artificial Orange Flower Water, Tri	2	cc.	Formula No. 1		
Orange Flower Water, Tri	ple 100	cc.	Lemon Oil	18	g.
Alcohol (90%)	900	cc.	Bergamot Oil	16	g.
223001101 (2-70)			Orange Oil, Sweet	5	g.
77			Lavender Oil, Extra	4	g.
Hungary Water			Mandarin Oil	3.2	g.
Rosemary Oil	20	cc.	Petitgrain Oil, Grasse	3.2	g.
Verveine Oil	7 -	cc.	Benzoin Resinoid	3.2	g.
Portugal Oil	1.5	cc.	Neroli Oil, Original Orange Oil, Bitter Lime Oil	2.8	ğ.
Limette Oil	1	cc.	Vrange Oil, Bitter	2.8 2.7	Ř.
Rosemary On Verveine Oil Portugal Oil Limette Oil Peppermint Oil Rose Water, Triple Alcohol (90%)	100	cc.	2311110 011	1	
Rose Water, Triple	100	ec.	Rosemary Oil	0.6	g.
Alcohol (90%)	800	ec.	Curvin Aldebyde (10%)	0.5	ū.
Let stand up to 6 months	before	mar-	Muscatal Sage Oil	0.3	σ.
keting.			Eugenol Cumin Aldehyde (10%) Muscatel Sage Oil Hysop Oil Culculum Oil	0.1	σ.
			Cardamom Oil	0.1	ø.
Eau de Lubin			Iris, Concrete (10%)	0.1	g.
Alcohol	650	cc.	Iris, Concrete (10%) Alcohol (96%)	1800	cc.
Portugal Oil		cc.	Water, Distilled	200	cc.
Neroli Oil	0.6	cc.	No. 2		
Jasmine, Absolute	0.0	cc.	Bergamot Oil	20	œ
Myrtle Oil Geranium Oil, French	3	cc.	Lemon Oil	14	g. g.
Geranium Oil, French	1.2	cc.	Lavender Oil		g.
Denion On			Benzoin Resinoid	5 5	g.
Bergamot Oil		cc.		5	g.
Civet Tincture	3	cc.			
Castoreum Tincture	_	cc.	Orange Oil, Sweet Mandarin Oil Petitgrain Oil, Paraguay Rosemary Oil Neroli Oil Muscatel Sage Oil	4	ø.
Peruvian Balm	3 6 24	cc.	Petitorain Oil, Paraguay	2.6	ğ.
Musk Tincture	o c	cc.	Rosemary Oil	2.3	₿g.
Tolu Balm Tincture	94	00.	Neroli Oil	2	g.
Benzoin Tincture	6	cc.	Muscatel Sage Oil	2	g.
Myrrii Iniciare	60	cc.	Jasmine Aldehyde	0.7	g.
Clove Tincture	00		Rosemary Oil Neroli Oil Muscatel Sage Oil Jasmine Aldehyde Resinoid Iris Alcohol (96%)	0.5	g.
			Alcohol (96%)		
Aqua Mellis			Alcohol (96%) Water, Distilled	200	cc.
Honey	5	g.	No. 3		
Bergamot Oil	8	čc.	Lemon Oil	20	g.
Lavender Oil, French	1	cc.	Heliotropin	7	g.
Clove Oil	1		Bergamot Oil, Natural	5	g.
Maga Oil	0.	5 cc.	Bergamot Oil, Artificial	6	g.
Coriander Oil	1	cc.	Terpinyl Acetate	4	g.
Sandal Wood Oil	1 3. 5 2	5 cc.	Neroli Oil, Artificial	4	g.
Benzoin Resinoid	5	cc.	Orange Oil, Sweet	4	g.
Sandal Wood Oil Benzoin Resinoid Musk Tincture (2%) Rose Water, Triple	2 100	cc.	Coumarin	2.4	5 g. 5 g.
Rose Water, Triple	100	cc.	Benzyl Acetate	1.	. B.

Chypre, Eau de Cologne Chypre, Eau de Cologne Chypre, Eau de Cologne Chypre, Eau de Cologne Chenon Oil 18 g. Bergamot Oil 16 g. Bergamot Oil 16 g. Bergamot Oil 12 cc. Chenon Oil 10 cc. C				
Ambre Eau de Cologne Bergamot Oil 20 g.	Ketone Musk	0.7 g.	Eau de Cologne for t	he Bath
Ambre Eau de Cologne Bergamot Oil 20 g.	Citral	0.6 g.	Bergamot Oil, Free of	
Ambre Eau de Cologne Bergamot Oil 20 g.	Alcohol (96%)	1600 cc.	Terpenes	17 cc.
Ambre Eau de Cologne Bergamot Oil 20 g.	Water, Distilled	400 cc.		
Ambre Eau de Cologne Bergamot Oil 20 g.		-	Terpenes	14 cc.
Bergamot Oil	Ambro Dr. J. A	O	Rosemary Oil	1.75 ec.
Namina	Paramet Oil	90 ~	Citrat	1.75 cc.
Namina	Lemon Oil	20 8.	Orango Flawer Water	340 cc.
Namina	Heliotropin	7 6.	Alcohol (96%)	1800 cc.
Namina	Ambrette Musk	2.6 2.	Water, Distilled	3600 cc.
Namina	Lavender Oil	2.6 g.		
Namina	Petitgrain Oil, Paragu	nay 2.6 g.	Tea Bay Bum	
Namina	Methyl Ionone	2.6 g.		
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Vanillin	2 g.	Manthal	16 g.
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Rose Oil, Artificial	2 g.	Glycerin C.P.	
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Nosell Od	0.7 g.	Glycerin (Soap Lye)	20 g.
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Coumarin	0.7 8	Rum Essence	80 g.
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Ambre, Artificial	0.6 %	Alcohol (96%)	2000 cc.
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Rose Absolute, Synthe	tic 0.1 g	Water, Distilled	800 cc.
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Alcohol (96%)	1800 cc.		
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.	Water, Distilled	200 cc.	Eau de Lavend	le
Chypre, Eau de Cologne Lemon Oil 18 g. Bergamot Oil 16 g. Rose Oil, Artificial 6 g. Lavender Oil 4 g. Coumarin 4 g. Sandal Wood Oil, East India 2 6 g. Ketone Musk 2.6 g. Muscatel Sage Oil, Artificial 2 g. Muscatel Sage Oil, Artificial 2 g. Bao-Eugenol 0.7 g. Patchouli Oil 0.7 g. Vanillin 0.5 g. Thyme Oil 0.5 g. Thyme Oil 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Mousse de Chêne, Absolute 0.5 g. Water, Distilled 200 cc. Eau de Cologne ''Russe'' Eau de Cologne ''Russe'' Eau de Cologne Perfume Bergamot Oil 12 cc. Ambre Infusion 12 cc. Bergamot Oil 12 cc. Phenyl Eth.1 Alcohol 0.6 cc. Vater, Distilled 300 cc. Vater, Distilled 300 cc. Vater, Distilled 200 cc. Bitter Almond Perfume Bitter Almond Oil 60 cc.			Lavender Oil, Barrême	
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume			(France)	40 cc.
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume	Chypre, Eau de	Cologne	Musk Infusion	12 cc.
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume	Lemon Oil	18 g.	Ambre Infusion	12 ec.
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume	Bergamot Oil	16 g.	Bergamot Oil	12 cc.
Rosemary Oil 2 g Muscatel Sage Oil, Artificial 2 g Huscatel Oil H	Rose Oil, Artificial	9 g.	Lemon Oil	6 ec.
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume	Coumaria	4 g.	Jasmine Aldebyde	2 cc.
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume	Sandal Wood Oil East	India 2 6 g.	Aladal (960)	1100
Rosemary Oil 2 g Museatel Sage Oil, Artificial 2 g Eau de Cologne Perfume	Ketone Musk	2.6 g.	Water Distilled	300 cc.
Australe Sage Oil, Artineial 2 K Eau de Cologne Perfume	Vetivert Oil, Java	2 g.	7711(1)	0.00 001
Australe Sage Oil, Artineial 2	Rosemary Oil	2 g.	Darfumas for Shaning	Crooms
Mousse de Chêne, Absolute 0 5 g. Alcohol (96%) 1800 cc. Water, Distilled 200 cc. Eau de Cologne "Russe" Bitter Almond Oil 60 cc. Bitter Almond Oil 60 cc.	Museatel Sage Oil, Art	ificial 2 g.		
Mousse de Chêne, Absolute 0 5 g. Alcohol (96%) 1800 cc. Water, Distilled 200 cc. Eau de Cologne "Russe" Bitter Almond Oil 60 cc. Bitter Almond Oil 60 cc.	Iso-Eugenol	07 g.		
Mousse de Chêne, Absolute 0 5 g. Alcohol (96%) 1800 cc. Water, Distilled 200 cc. Eau de Cologne "Russe" Bitter Almond Oil 60 cc. Bitter Almond Oil 60 cc.	Patchouli Oil	0.7 g.		100 g.
Mousse de Chêne, Absolute 0 5 g. Alcohol (96%) 1800 cc. Water, Distilled 200 cc. Eau de Cologne "Russe" Bitter Almond Oil 60 cc. Bitter Almond Oil 60 cc.	Vanillin Nordi Oil	0.5 g.	Lemon Oil	50 g.
Mousse de Chêne, Absolute 0 5 g. Alcohol (96%) 1800 cc. Water, Distilled 200 cc. Eau de Cologne "Russe" Bitter Almond Oil 60 cc. Bitter Almond Oil 60 cc.	Thyma Oil	0.5 g.	Portugal Oil	35 g.
Eau de Cologne "Russe" Bitter Almond Perfume Bitter Almond Oil 60 cc.	Mousse de Chêne Abso	olute ()5 g. l	Layendar Oil	30 g
Eau de Cologne "Russe" Bitter Almond Perfume Bitter Almond Oil 60 cc.	Alcohol (96%)	1800 cc.	Petitorain Oil	30 %.
Eau de Cologne "Russe" Bitter Almond Perfume Bitter Almond Oil 60 cc.	Water, Distilled	200 cc.	Neroli, Synthetic	20 g.
Eau de Cologne Russe Bitter Almond Oil 60 cc.	•		, ,	·
Eau de Cologne Russe Bitter Almond Oil 60 cc.			Bitter Almond Pc	rfume
Lawender Oil 9 6 Lawender Oil 10 cc.	Eau de Cologne '			
Bergamot Oil 9 K Lavender Oil 5 cc.	Lemon Oil	9 g.	Bergamot Oil	10 cc.
Methyl Ionone 0 g G Helhotropin 4 g Fancy Perfume	Bergamot Oil	9 g.	Lavender Oil	5 cc.
Helotropin	Methyl Ionone	6 g.	Approximation of the second se	
Lavender Oil 150 cc.	Heliotropin	4 g.	Fancy Perfum	e
NoRagemon 2.6 g. Portugal Oil 450 cc.	Lavender On	3 g.		
Retone Musk 2 g. Bergamot Oil, Synthetic 750 cc.	Vanillin	2.6 g.		
Rosemary Oil 2 g Lenon Oil 150 cc.	Ketone Musk	2 g.	Bergamot Oil, Synthetic	750 cc.
Linalyl Åcetate 2 g. Ambrette, Musk 0.7 g. Neroli Oil 0.7 g. Almond Perfume Counarin 0.6 g. Ambre, Artificial 0.6 g. Alcohol (96%) 1800 cc. Hehotropin 125 g. Water, Distilled 200 cc. Musk, Tincture 50 g.	Rosemary Oil	2 g.	Lemon Oil	150 cc.
Ambrette, Musk 0.7 g. Neroli Oil 0.7 g. O.7 g. Ounnarin Almond Perfume Counnarin 0.6 g. Ambre, Artificial 0.6 g. O.7 g. Ounnarin Peru, Balsam 100 g. Ounnarin Alcohol (96%) 1800 cc. Heliotropin 125 g. Ounnarin Musk, Tincture 50 g. Ounnarin	Linalyl Acetate	2 g.	Benzaldehyde	30 cc.
Neroli Oil	Ambrette, Musk	0.7 g.	*	
Countarin 0.0 g. 1.0 g.	Neroli Oil	0.7 g.	Almond Perfun	ne
Alcohol (96%) 1800 cc. Heliotropin 125 g. Water, Distilled 200 cc. Musk, Tincture 50 g.	Coumarin	0.6 g.		
Water, Distilled 200 cc. Musk, Tincture 50 g.	Ambre, Artificial	1800 cc	Heliotropin	125 g
Travel, Diovines	Water Distilled	200 cc.	Musk, Tincture	50 g.
	Tracel, Distinct			

Vanillin	15 g.	Perfume for Choleste	erin C	ream	8
Almond Oil	10 g.	1. Orange Flower Wa			
Neroli, Synthetic	5 g.	water:			
Lavender Perfume		Neroli Oil, Artificial		9	g.
Lavender	75 g.	Aubépine		1	g.
Lavender Spike Oil	75 g.	2. Rose Water instea	d of		tilled
Geranium Oil	75 g.	water:			
Commarin	and the Bear	Rose Oil		1	g.
Sandal Wood Oil	**************************************	- Géranium Oil, African		1	g.
Bergamot Oil	100 g.	Bergamot Oıl		5	g.
Lemon Oil	25 g.	3. Rose Water instea	d of		tilled
Rose Perfume		water:			
Pelargol	100 g.	Geranium Oil		5	g.
Diphenyl Oxide (1:1)	25 g.	Anisaldehyde		5	g.
Vanillin	10 g.	Linalylacetate		2	g.
Geraniol	75 g.	Eugenol		1	g.
Terpineol	20 g.	The three mixtures	0.00		
Violet Perfume		creams made with Rose V			
Bergamot Oil	100 g.	Flower Water instead of			
Iris Resmoid	30 g.	(Usual percentage of per			
Neroli	25 g.	` • • •		′	
Benzoin Infusion	75 g.				
Terpineol	50 g.				
Violet (5187, Heine)	125 g.	PERFUME BA	ASES		
Jasmine Flower Oil	40 g.			,	
Fixol-Violet	50 g.		Ę	,	1
Extract, Rose			Mown	Chypre	يدا
Red Rose Flower Oil	40 cc.		New Hay	Į ā.	Locust
Nerol	30 cc.		98	Ę,	١٤
Phenyl Ethyl Alcohol	20 cc.				
Jasmine Aldehyde	16 cc.	Alpha Ionone	10		_
Neroli Oil	12 cc.	Citrouellol	20		
Ambrette Musk	10 cc.	Amyl Saheylate	100	25	5.5
Rose Absolute, Synthetic	9 cc.	Anisic Aldehyde	20	-	
Iris, Concrete	5 ec. 3 ec.	Countarin	5		-
Tuberose, Artificial Bergamot Oil	2 ec.	Vanillin	5	5	
Norganiot (71 Norgango Artificial	2 cc.	Heliotropin	7 10	10	7
Narcisse, Artificial Vetivert Oil, Java	1 cc.	Lanolool Petitgrain	10	10 20	2.5
Sandal Wood Oil, East Indi		Jasmine, Artificial	20	25	4.5
Alcohol (96%)	1500 ec.	Patchouli Oil	ĩ	25	1
Water	150 сс.	Aldehyde C10, 50%	ī	_	.15
T 11 12 A		Iso Eugenol	5	_	2.3
Lilac Perfume	• •	Phenyl Ethyl Alcohol	-	25	20
Anisic Aldehyde	10 cc.	Musk Xylol	-	25	
Jasmine, Synthetic Heliotropin	10 cc. 5 cc.	Coparba, Balsam	-	15	-
Phenyl Ethyl Alcohol	5 cc.	Birch Tar Lemon Oil	-	10	-
Phenyl Éthyl Alcohol Phenyl Acetaldehydo	5 cc.	Bergamot Oil	-	3 100	_
Oil Bergamot	3 cc.	Rose, Artificial		75	
Musk Ketone	3 cc.	Cedar Oil		15	
Styrax Resin	2 cc.	Phenyl Acetic Aldehyde,			
Oil Ylang Ylang	2 cc.	50%			1
Terpineol	55 cc.	Phenyl Acetic Acid	-		.25
Individual touches may be i		Hydroxycitronellol	-	_	12.5
the above by the sparing use	of any or	Cinnamic Alcohol	-		3.5
all of the following: amyl		Canauga Oil	-	_	3
acetophenone, methyl anthrani	ute, penzyl	ate, 5%		_	1.0
acetate, cinnamic alcohol, b		Geranyl Acetate			1.3
meg.	_ VII MUV	Amyl Cinnamic Aldehyde			5
·		1			

				4		1	_	l m
	Flowery Bouquet	Bouquet	Oriental	Oriental		Flowery Bouquet	Bouquet A	Bouquet B
	Mo Mo	bno	ıen	ien		g a	9	nb
	Eğ	Ä	ō	Ö		30.0		30 n
					A CONTRACTOR OF THE PARTY OF TH	-		
Rose Geranium Oil	100				Methyl Phenyl Acetate		40	-
Rose, Artificial	20		350	170	Musk Ambrette		100 50	
Valley Lily, Artificial	500 200	500	350a 110	100	Para Cresyl Phenylacetute Vanillin		30	
Terpineol	200				Aldehyde C., 5%		100	100
Bois de Rose	200			-	Olibanum Gum, 2:1		150	
Coumarin	30	100	-		Terpincol	-:		200
Amsic Aldehyde	20 150	20	30	30	Hydroxyertionellal Cananga Oil	1		100
Methyl Anthranilate. Civet Tincture	50		100	60	Rose Geranium Oil	1		100
Hyacinth, Artificial.	100				Coumarm			30
Benzyl Benzoate	200	200	200	100	Amsic Aldeliyde			20
Musk Ambrette	50		50	30	Methyl Authramlate			100
Openax	200		200 100	100 50	Civet Tineture Labdanum			100
Oak Moss, Liquid Cananga Oil	100				Corrander Oil			20
Lavender Oil		20	20	10	Castoreum, 10% Ambergris Tincture			100
Bergamot Oil		100			Ambergris Tineture			100
Cassia Oil		10	-	-				
Tuberose, Artificial . Methyl Heptine Car-		100		_			07	
bonate, 5%	-	100		_		4	t	et]
Geraniol		100	-			pre	ğ	n _b
Vanillin		100			!	Chypre A	Bouquet C	Bouquet D
Musk Ketone	-	50		-		0	m	щ
Orange Blossom, Ar- tificial	_		610	100		004		
Jasmine, Artificial			110		Jasmine, Artificial	200 400	500 200	80 500
Vetivert Oil				100	Oak Moss, Liquid	500	100	
Jasmine Aldehyde	-			200		1000	-	
Petitgrain Oil Phenyl Ethyl Alcohol		-		30	Rose, Absolute	400		
Linalyl Acetate				50	Patchouli Oil	500) 200		200
Linalool				50	Vanillin	100		
					Coumarin	200		
		ر ر		آھ ا	Indol, 5%	100		
		7.E	4	Rouquet B	Hydroxycitronellal Lemon Oil, Terpencless.	200 30		
		5 2	ant.	ň	Phenyl Ethyl Alcohol		100	
		Flowery Bouquet	Bouquet	ğ	Methyl Ionone		500	
		4,2	æ	E C	Aldehyde C ₉ , 50%		40	
		-			Methyl Heptine Carbon-		50	
Aldehyde C	• • • •	20	20	20	ate, 10%		200	
Oak Moss, Liquid		100 500	200	200	Iso Butyl Salicylate		200	
Jasmine Liquid, Abs. Rose, Artificial		500	1000		Rhodinol		500	150
Iso Butyl Sahcylate .		200	100	100	Lilac, Artificial		500	
Methyl Ionone		500	-	300	Valley Lily, Artificial Bois de Rose		500 200	
Lilac, Artificial	• • • •	500	200	300 200	Cassie, Artificial			60
Musk Ketone Methyl Heptine Carbon		200	_	1500	Benzyl Benzoate		_	1000
5%		50	-		Dicthyl Anthramlate			50
Valley Lily, Artificial		500	200		Benzyl Acetate	_		300
Bois de Rose		200	200	200	Tolu, Balsam		_	300
Melittis (Givaudan) Orange Blossom, Arti		200		300	Rose, Artificial	***		90
Orange Proposing Hitt								

Oak Moss, Liquid Bergamot Oil, Terpencless. Linalyl Acetato Sweet Orange Oil Valley Lily, Artificial Narcissus Absolute Jasmine, Artificial Rhodinol Alcohol Co	200 150 200 300 100 400 200 70	0	Raldeine D Lemon Oil Rhodinol Alpha Ionone Hydroxyeitronellal Cananga Oil Aldehyde C ₁₂ , 5% Methyl Heptin Carbonate 10%		20 1 1	100
Aldehyde Co, 5% Linalool Geranyl Acetate Methyl Phenylacetate Alpha Ionono Vetivert Oil Terpincol Coumarin Vanillin Musk Ketone Canada Snake Root Oil	100 200 200 50 100 100 200 100 100 100	30 100 100	Cassie, Artificial		=	100 300 100 175
Hydroxycitronellal Geraniol Phenyl Acetic Aldehyde, 50% Phenyl Ethyl Alcohol Anisic Aldehyde Rose, Artificial Labdanum	= = = = = = = = = = = = = = = = = = = =	2000 50 50 300 20 30 100	Bergamot Oil Bois de Rose Benzyl Alcohol	1500 150 150 150	S Sweet Pea	8000 Heavy
	Щ	1	Phenyl Ethyl Alcohol Indol, 5% Hydroxyettronellal Orange Blossom, Artifical Cananga Oil Jasmine Absolute	300 50 250 250 150 300	200 200 200 200	200 - 200
	Bouquet 1	Violet	Amyl Cinnamic Aldehyde Benzylidene Acetone . Heliotropin	100	150 450 50	100 —
Bergamot Oil, Terpeneless. Linalyl Acetate Jasmine, Artificial Aldehyde Co, 5% Vetivert Counnarin Rose Geranium Oil Rose, Artificial Bny Oil, Terpeneless Eugenol Pettigrain Oil Bergamot Oil Indol, 5% Ambreol Lavender	200 100 500 100 100 400 200 100 300 100 400 300 150	= = = = = = = = = = = = = = = = = = = =	50% Terpineol Iso Butyl Phenylacetate Rose, Artificial Tolu Alcohol Co Benzyl Benzoate Anisic Aldehyde Lavender Oil Tolyl Acetate Vanillin Oak Moss, Liquid Aldehyde C10, 5% Diethyl Anthranilate Ambreol		100 1000 120 80 150 60 150 — — — —	

				To the second se		
		Carnation	Honeysuckle	Lilac	Orange Blossom	Heary Modern Oriental
Eugenol Jasmine, Artificial Heliotropin Rose, Artificial Phenyl Ethyl Alcohol Orange Blossom, Artific Ocallet Orris Liquid, 10% Musk Ketone Ambreol Benzyl Iso Eugenol Bergamot Orl Indol, 5% Hydroxycatronellal Benzyl Acetate	ual.	1600 400 400 100 50 100 150 100 100 	2000 2000 2000 	Linalyl Acetate	100 15 10 -	50 30 10 6 5 3
Benzyl Butyrate Benzyl Formate Benzyl Fropionate Benzyl Benzoate Bois de Rose Aurania Cananga Oil Amyl Cinnamic Aldehyde Para Cresol, 10% Petitgrain Oil	e	-	500 200 2000 2000 700 800 1000 500 100	Lily-of the Valley Flower Geraniol, from Palmarosa Oil Linaloof, from Rosewood Oil Phenylacthyl Alcohol Phenylacetaldehyde Dimethyl a ctal a lonone Benzaldehyde Jasmine Flower Oil, Arti- ficial Rose Oil, Artificial,	25 12.5 15 5 1.5 0.1	g. g.
Citronellol	10 20	- 5 5 -	3 -	Extra Fine Lake Flower Oil, Artificial Ylang Ylang Oil, Manila Rhodinol Corander Oil, Terpene Free Hydroxyeitronellal Dimethyl- acetal Hydroxyeitronellal Diethyl- acetal	8 25 4 10 0,5 20 40	g. g
hyde Methyl Acetophenone Hydroxycitronellal Phenyl Ethyl Alcohol Linalool Terpineol Methyl Para Cresol Musk Ketone	10 -2 11 22 -2 1 -	5 1 5 1 5 1 5 1 5 1 7 7 7 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	50	Lilac Flower Oil Ylang Ylang Oil, Mamla Jasmine Flower Oil, Artificial Rhodinol Acacia Flower Oil, Artificial Hydroxyettronellal Diethyl- nectal Terpincol, Extra Phenylacetaldehyde Dimethylacetal Aubépine (from Anethol) Hehotropin Iso Eugenol Vanillin Octyl Acetate (10%) in Benzyl Alcohol	30 20 4 2 12 1.5 0.5	

Perfume Oil, Type "Te	osca''	
Formula No. 1		
Orange Oil, Sweet,	0 5	
Calabrian	8.5	cc.
Bergamot Oil, Extra Fine,	17	
Reggio	17	cc.
Lemon Oil	19	cc.
Ylang Ylang, Genuine	6	cc.
Rose Oil, Genuine,		
Bulgarian	2.5	cc.
Jasmine, Pure	1.3	cc.
Coumarin	6.5	g.
Musk, Artificial,	_	
"Ambrette"	1	g.
Musk, Artificial,	_	
"Ketone"	1	g.
Cedar Wood Oil, Rectified	5.5	cc.
Neroli Oil, Genuine Geraniol, C.P.	2.5	ec.
Geraniol, C.P.	4	ec.
Phenylethyl Alcohol	1.5	ee.
Benzoin Extract, Filtered	5	ec.
Petitgrain Oil	1.5	ec.
Linaloe Oil, Cayenne	6	cc.
Linaloe Oil, Cayenne Sandal Wood Oil,		
East Indian	5.5	ee.
Indol (100%)	0.07	ce.
Iris Oil, Genuine, Concrete	1.5	ee.
Castoreum (100%)	0.05	g.
Busilicum Oil	0.03	će.
Undecyl Aldehyde (100%)	0.05	cc.
Undecyl Aldehyde (100%) Mousse de Chêne, Laquid	0.5	ce.
Vanillin	3	g.
Menthol	0.5	g.
No. 2		
Bergamot Oil, Extra Fine	11	cc.
Lemon Oil	26.5	ec.
Orange Flower Water Oil,	20.0	
Genuine Water On,	1	cc.
Ylang Ylang Oil, Genuine	9	cc.
Sandal Wood Oil		
Sandal Wood Oil, East Indian	8	cc.
Amyl Salicylate	3.5	cc.
Iris Oil, Genuine, Concrete	1	cc.
Civet, Genuine (100%)	0.22	ce.
Patchouli Oil	1.5	cc.
Coumarin	4	g.
Vanillin	5	
	3.5	g.
Rose Oil, Bulgarian	1.5	cc.
Petitgrain Oil Musk Artificial	1.0	
Musk, Artificial,	6	œ
Germiol C.P	6.5	g.
Gernniol, C.P. Benzoin Extract, Filtered	5	cc.
Underel Aldehyda (100c/.)	0.2	
Undecyl Aldehyde (100%) Birch Tar Oil,	0.4	g.
Twice Rectified	0.03	cc.
Coder Wood Oil Rectified	2	cc.
Neroli Oil, Genuine	0.5	cc.
Linaloë Oil, Cayenne	2	cc.
Oponopor Extract	0.05	cc.
Opoponax Extract	2	
Jasmine Oil, Pure	_	ce.
The above-mentioned perfu	me cor	nposi-
tions should be made up 1-29	% in 8	90%

pure alcohol and kept in the dark, shaking from time to time, and filtering after a few weeks.

Perfume Oil, Type "Quelqu Tart ("Herb") Ty		eurs'
Formula No. 1	Γ-	
Olibanum Oil	3	cc.
Geraniol, C.P.	7.5	cc.
Alpha Amyl Cinnamic	7.0	ш.
Aldeliyde	2.36	cc.
Citral	5	cc.
Geranium Oil, Réunion	3.5	cc.
Benzyl Alcohol	10	cc.
Linalyl Acetate	7	cc.
Hydroxycitronellal, C.P.		
(100%)	14	cc.
Heliotropin, Crystallized Cananga Oil, Java	10	g.
	13	cc.
Ionone for Soaps	4	cc.
Methylnonyl Acetaldehyde		
(100%)	0.14	cc.
Benzyl Acctate, Free of		
Chlorine	6	cc.
Linaloë Oil, Cayenne	3	ec.
Terpineol, C.P.	11	cc.
Musk, ''Ambrette,''		
Artificial	0.5	g.
No. 2		
Benzoin, Extract	3	cc.
Olibanum Oil	1.36	cc.
Citronella Oil, Colombo	3	ee.
Cananga Oil, Java	10	ec.
Hehotropin, Crystallized	6	\mathbf{g} .
Linaloë Oil, Cayenne	7	cc.
Hydroxycitronellal, C.P.	-	
(100%)	7	cc.
Benzyl Acetate, Chlorine-	3	
Free	$\frac{3}{26.5}$	cc.
Terpincol, C.P. Citral	3	('C.
Methylnonyl Acetaldehyde	J	cc.
(100%)	0.14	cc.
Geranium Oil, Réunion	5.5	cc.
Ionone for Soaps	5.5	cc.
Phenylethyl Alcohol	5	ec.
Linalyl Acetate	4.5	cc.
Anise Aldehydo	6.5	cc.
Alpha Amyleinnamic	0.0	30.
Aldehyde	3	cc.

Perfume Oil, Type "Quelques Fleurs"
For Fine Soaps (Soft Type)
Cananga Oil, Java 9 cc.
Benzyl Acctate, Free of
Chlorine 5 cc.
Linalyl Acetate 5 cc.
Linaloë Oil, Cayenne 2.3 cc.
Heliotropin, Crystallized 8 g.
Geraniol, C.P. 8 cc.

	JOHN THE L	AND DRUGO
Musk, "Ambrette,"		C
Artificial	95 ~	Cutic
	3.5 g.	Glycerol
Bergamot Oil	2 cc.	Potassium Hyd
Phenylethyl Alcohol	3.5 cc.	Water
Benzyl Alcohol Alpha Amylcinnamic	9 cc.	Perfume
Alpha Amylcinnamic		Basic Red Dye
Aldehyde	0.5 cc.	
Terpineol, C.P.	21 cc.	The potassium
Indol, Crystallized	0.06 g.	in the water a
Laura Od Consino		added. The per
Lemon Oil, Genuine		terpeneless lemor
Anise Aldehyde	4 cc.	is added to give
Hydroxycitronellal, C.P.	9 cc.	bottle.
Methylnonyl Acetaldehyde		Dott It.
(100%)	0.14 cc.	1
` , , ,		1
grand and an artist of the second		Cutic
D . C O'l - ((()) I	3.4	For
Perfume Oils "Chypre I	extract.	1
Formula No. 1		Light Turbine
	33 ec.	to suit.
Bergamot Oil		1
Geranium Od, Réunion	2 - cc.	Diglycol Laura
Rose Oil, Genuine, Bulgari	an 3.5 cc.	
Ylang Ylang Oil, Genuine	25 cc.	Deodorized Ke
Rosemary Oil	4 cc.	Perfume
Coumarin	8 g.	l .
Lavender Oil, Genuine	6 cc.	Ohve Oil
Jasmine, C.P.	2.4 cc.	Petroleum Jell
37		Red Dye Oil S
Vanillin		nea Dye On B
Anise Aldehyde	5.5 cc.	
Cedar Wood Oil, Rectified	1.5 cc.	Perfune Lilac,
Patchouli Oil, Genuine	0.5 cc.	A lower price
Mousse de Chêne, Decoloria	ted 3 - cc.	pared by using
Opoponax Extract	2 ec.	mineral oil. The
Linaloë Oıl, Cayenne	18 cc.	
Civot Genuine (100%)	0.6 %.	be nearly white.
Civet, Genuine (100%) Musk, "Ambrette," Artific	dal 45 g	a low heat and
Musk, Ambrette, Artine	ш то Б.	The dye is masce
No. 2		tion of the oil a
Lemon Oil	12 cc.	tout the entire
Bergamot Oil	9 ec.	added in amor
Benzyl Acetate, Free from		strength of the p
		I was a second
Chlorine		
Cedar Wood Oil, Rectified	9.5 cc.	N
Benzyl Benzoate	6 cc.	1
Hydroxycitronellal, Pure		For
(100%)	5 cc.	Amyl Acetate
Geraniol, C.P.	7 cc.	
Vanillin	4 g.	Methyl Alcoho
Dannin Franci Lultored	5.5 6.	Nitrocellulose
Benzoin Extract, Filtered Sandal Wood Oil, East Indian	0.0 ***	Benzoin
Sandai Wood Oil,	F 43	Carmoisine (19
East Indian	5 cc.	Solution)
Geranium Oil, Réunion	3 ec.	1
Couma rin	2 g.	1
Rose Oil. Genuine, Bulgari	ian 1 cc.	D. t. I. A . et . A .
Lingloë Oil, Cavenne	2.5 cc.	Butyl Acetate
Rose Oil, Genuine, Bulgari Linaloë Oil, Cayenne Musk, "Ambrette,"		Ethyl Acetate Ethyl Alcohol
Artificial	1.5 g.	Ethyl Alcohol
Artificial	1.5 cc.	Butyl Alcohol
Patchouli Oil, Genuine	3 cc.	Damar
Labdanum Extract		Color
Civet, Genuine	0.3 g.	1 ~~~~
	0.7 cc.	1
Olibanum Extract		Methyl Ethyl
Olibanum Extract Iris Oil, Genuine, Concrete	1 cc.	
Iris Oil, Genuine, Concrete Mousse de Chêne, Decolori	zed 2 cc.	Resorcinol Dis
Iris Oil, Genuine, Concrete Mousse de Chêne, Decolori	zed 2 cc. 5 cc.	Resorcinol Dis Ethyl Lactate
Olibanum Extract Iris Oil, Genuine, Concrete Mousse de Chêne, Decolori Ylang Ylang Oil, Genuine Phenylethyl Alcohol	zed 2 cc. 5 cc. 5.5 cc.	Resorcinol Dis

Cuticle Remover		
Glycerol	20	OZ.
Potassium Hydroxide	4	oz,
Water	76	oz.
Perfume	0.3	oz.
Basic Red Dye	t	race

hydroxide is dissolved n nyaroxide is dissolved and the glycerol then ifume usually used is no in oil. Just enough dye same a pink color in the

cle Softene<mark>r</mark>

mula No. 1

Oil - color and perfume

No. 2			
Diglycol Laurate	10	oz.	
Deodorized Kerosene	10	oz.	
Perfume	to	to smt	
No. 3			
Olive Oil	88	OZ.	
Petroleum Jelly	12	OZ.	
Red Dye Oil Soluble			
to a mini	e anlor t	PD 00	

to a pink color trace, enough, about 0.3 oz.

d product may be pre-a medium bodied white a medium bodied white the petroleum jelly should. This jelly is melted at added to the olive oil. The perfunction of this paste is used to mass. The perfune is unit varying with the particular product used.

ail Polish rmula No. 1

700 g.

300 g. 50 g. 100 g. % Alcoholic 50 cc. or to suit

No. 2	
Butyl Acetate	250 g.
Ethyl Acetate	150 g.
Ethyl Alcohol	400 g.
Butyl Alcohol	200 g.
Damar	5 g.
Color	to suit

No. 3	
Methyl Ethyl Ketone	650 g.
Resorcinol Diacetate	100 g.
Ethyl Lactate	200 g.
Nitrocellulose	100 g.

Sandarac		5 g	ζ.
Color		to sui	t
~	 		-: 1

Sometimes the polish is perfumed with a little ionone or ylang ylang oil, but more often this is not done.

No. 4

Nitrocellulose	
Viscosity)	225 g.
Damar	75 g.
Butyl Acetate	25 g.
Butyl Alcohol	20 g.
Ethyl Acetate	15 g.
Alcohol	40 g.
Carmine Red	sufficient to color

Nail Polish Powder

Putty Powder (Tin Oxide)	40 oz.		
Infusorial Earth (325 Mesh)	55 oz.		
Stearic Acid (Powdered)	5 oz.		
Color (Pigment)	to suit		
Perfume	to suit		

Removers, Nail Polish Formula No. 1

The nail polish remover consists chiefly of the solvent alone. It has been found, however, that butyl stearate has a particularly rapid action on the film, and many modern removers make use of it in conjunction with other solvents. An effective remover can be made by mixing butyl stearate 1 part, amyl acetate 3 parts, and acetone 4 parts. Diglycol laurate is also included to prevent brittleness of nails (about 1-2%).

No. 2	
Amyl Acetate	1 oz.
Acetone	1 oz.
No. 3	
Amyl Acetate	1 oz.
Alcohol	1 oz.
Acetone	1 oz.
Diglycol Laurate	1/8 oz.

Eyebrow Pencils

Apart from those methods which serve to preserve the eye region in good physical condition, actual beauty treatment is now practiced on a very considerable scale. Coloring of the eyebrows, painting of the eyelashes and shading of the eyelids are now important components of face cosmetics, the greatest attention being devoted to the first operation. Coloring of the eyebrows or their simulation after complete shaving is effected with colored wax pencils. As already mentioned, ordinary pure charcoal pencils tend to cause falling-out and drying of the hair.

Ingredients used in preparing the wax pencils are white wax, benzoated tallow, cocoa butter, petroleum oil and olive oil. The pigments are lamp black, umber, and ochre. Large manufacturers find it economical to use pigment grinding ma-chines and other equipment of the most modern design, but small concerns can nevertheless cope with the production of these cosmetics. The base comprises a composition made up from 110 g. fine petroleum oil, 60 g. white ceresine, 15 g. white was 240 g. benzoated tallow, and 1 g. coumarin. The fatty base is thor-oughly ground with the pigments, the molten base being gradually stirred into the very finely powdered pigment contained in a mortar. After thorough trituration the mixture is again warmed, digested for about half an hour on a water bath, and again allowed to cool. As soon as the mass begins to thicken, it is again vigorously stirred and forced through a fine-mesh sieve by applying powerful pressure with the pestle. Lumps and impurities are retained upon the sieve. The preparation which passes through the mesh is then again thoroughly mixed, with gentle heating before casting. The mass should be neither too hot nor too fluid when being cast, since settlement of the insoluble pigment will result in lack of uniform coloration. Oilsoluble dyestuffs will certainly only enter into consideration in exceptional cases. According to another process, the melt is prepared from 2 parts cocoa butter, 2 parts ceresine, and 1 part olive oil. Into this is stirred 0.6 part dyestuffs (i.e., about 10% of the total gross weight), which has previously been ground up with a little olive oil.

As soon as the mass has reached the state when it can just be east, it is emptied into metal moulds. As a rule these impart the required taper to the pencils, but if this is not the case they are tapered after removing from the moulds and wrapped in thick metal foil while leaving the points exposed.

Eyelid Pencils

The production of shad of tones on eyelids can be effected with pencils, the composition of which is very similar to that of the eyelrow pencils. The mass consists of the fatty base detailed above with the addition of about 20% ceresine. The color scale is somewhat more varied in the case of these pencils, since a wider range of tones can be induced in the usual brown and bluish black shades. Chestnut is obtained by mixing 225 g.

pale umber and 150 g. mahogany brown with 1000 g. of the molten wax mass.
For dark brown tones mix with the same
quantity of wax 300 g. of a brun foncé;
black shades require for the same wax
quantity 100 g. zinc white, 120 g. ultra-
marine, and 4 g. lamp black.
Regarding the perfuming of those

preparations, these should generally be of a very refined character. About 5 to 10 g. of perfume are required for each kilogram of mass. In cases where a fancy perfume is desired, preference should be given to one with a fresh natural odor.

Brown Eyebrow	Pencil	
Burnt Sienna	80 g	
Burnt Umber	100 g	
Hard Paraffin	420 g	
Soft Paraffin, Yellow	400 g	

Eyebrow and Eyelash Softener Formula No. 1 Castor Oil 07. Almond Oil 60 0Z. ¾ ∪z. Perfume No. 2 Diglycol Laurate 100 oz. Acetic Acid, Glacial 1/4 oz. 200 Mineral Oil, Medicinal oz. No. 3 200 g. Beeswax 300 g. Cocoa Butter Melt together and add: Peanut Oil 750 g.

Lipsticks (and Eyebrow	Pencils)
Paraffin	2 oz.
Vaseline Oil, White	3 oz.
Beeswax, White	1 oz.
Ozokerite Ceresine	3 oz.

2 g.

1 υZ.

Colors: For 100 parts use:

Moldex or Other Good

Preservative

Titanium Dioxide

Fixation Red (Fixierrot)
I No. 6
Medium Ted (Mittelrot) 3.5 oz. No. 28 22 oz.

Other red ves used: Carmine, Naka-rat, Fixierrot, Therry Red, Orient Red.

After Shave Lotions Formula No. 1 Glycerin 2 g. Lactic, Citric, or Phos-0.2 g. phoric Acid Menthol 0.5 g.

Alum	0.3 g.
Perfume	0.5 g.
Alcohol (45%)	96.5 g.
No. 2	
Glycerin	5 g.
Alum	1 g.
Zine Sulphophenolate	0.5 g.
Propyl Alcohol, C.P.	10 g.
Rose Water	10 g.
Perfune	0.5 g.
Alcohol (45%)	72.5 g.
No. 3	
Alcohol (40%)	1000 cc.
Glycerin, C.P.	
Aluminum Lactate	
Citric Acid	
No. 4	2 g.
Zinc Sulyhophenolate	0.5 g.
Alcohol (96%)	15 cc.
Witch Hazel	10 g.
Peruvian Balm	0.25 g.
Glycerin, C.P.	1 g.
No. 5	
Distilled Water	20 cc.
Isopropyl Alcohol	4 cc.
Alcohol	4 cc.
Alum	1 g.
Glycerin	0.5 cc.
Zinc Sulphophenolate	0.25 g.
No. 6 (Cloudy)	
Emulsone B	50 g.
Borie Acid	50 g.
Isopropyl Alcohol	100 g.
Diethylene Glycol	200 g.
Titanium Dioxide	
Distilled Water	60 g. 4 l.
Menthol	2 g.
Moldex or Other Preservat	
	-

Shaving Creams, Foaming

Formula No. 1	
Stearin	25 g.
a. Stearin Coconut Oil Fatty Acid	8 g.
b. Caustic Potash (50° Bé.) Water Glycerin	15 g.
b. { Water	50 cc.
COlycerin	4 g.
o. Stearin	3 g.

Melt up a, then introduce the solution b with stirring. Stir until cooled, then introduce c. When homogeneous, cover container and let stand over night. Perfume is added the next morning, optionally together with alcohol. Keep 8-14 days in earthenware jars, stir with a wooden rod on each day. In this time, the cream should become softer. If not, treat with a little caustic potash solution (20° Bé).

Perfume: Lavender, Rose, Violet,

THE C	CITEMICA	D PORMODARI	
Benzaldehyde, or with Eau de Cologne or Chypre.		No. 4	80 g.
No. 2		LOTing Off	100 g.
Palm Oil Fatty Acid,		Tallow	75 g.
Bleached	25 g.	Coconut Oil	60 g.
Olive Oil Fatty Acid	25 g.	(Caustic Potash (38° Bé.)	160 g.
Coconut Oil Fatty Acid	10 g.	b. Caustic Potash (38° Bé.) Glycerin Water	25 g.
Water	35 cc.	Water	15 g.
Caustic Potash (50° Bé.)	25 g.	c. Stearin	10 g.
Method as in No. 1. No. 3		As in No. 1.	
Stearin	30 g.		
Coconut Oil, or Fatty Acid	15 g.	D11 01 . 1 . 0	
Olive Oil, or Fatty Acid	10 g.	Brushless Shaving Crea	
Caustic Potash (28° B6.)	27 g.	1. Glycosterin	25 oz.
Water	32 cc.		10 oz.
Glycerin	6 g.	Peanut Oil Water	5 oz. 60 oz.
Stearin	3 g.	Moldex or Other Good	60 oz.
Method as in No. 1.		Preservative	0.2 oz.
No. 4		2. Stearic Acid	
Stearin	30 g.	Olive Oil	20 oz. 6 oz.
Coconut Oil	11 g.	Lanolin	2 oz.
Caustic Potash (50° B6.)	17 g.	Glycerin	6 oz.
Water	30 cc.	Triethanolamine	2 oz.
Glycerin Turkey Red Oil (100%)	10 g. 2 g.	Sodium Carbonate	1 oz.
to neutraliz	o alkali		63 oz.
***************************************		Perfume	to suit
Shaving Cream, Foamin Formula No. 1	ng .	-	
	20 ~	Soapless Shaving Prepara	tions
a. Stearin Peanut Oil, or Fatty Acid	30 g. l 10 g.		
Coconut Oil, or Fatty Acid	1 14 g.	German Patent 604,77	*
(Caustic Potash (38° Bé.)	28 g.	Formula No. 1	
b. Caustic Potash (38° Bé.) Water	20 g.	Glycol Stearate	100 g.
Glycerin (28° B6.)	12 g.	Water	400 g.
c. Stearin	5 g.	No. 2	
Mix a in the order of their		Absorption Base (Parachol)	
points (lowest first), melt up	to 60-70°	White Beeswax	25 g.
C., then stir in b, warm to 65	° C. Stir	Water	100 g.
until cool, add o (melted),	stir thor-	No. 3	
C., then stir in b, warm to 65 until cool, add c (melted), oughly, let stand over night. N	ext morn-	Glycol Palmitate	100 g.
ing stir up thoroughly, adding	pertume.	Petrolatum Water	100 g.
Cover, let stand, and fill into ea	rthenware	No. 4	200 g.
jars on next day.			100
No. 2		Diglycol Laurate Lanolin	100 g.
Bleached Palm Oil Fatty Ac	id 50 g.	Petrolatum	100 g. 50 g.
Olive Oil Fatty Acid	50 g.	Water	100 g.
Coconut Oil Fatty Acid	20 g.	No. 5	-04 P.
Water	70 g.	Stearic Anilide	100 g.
Caustic Potash (50° Bé.)	50 g.	Glycol Stearate	300 g.
Method as in No. 1.		Absorption Base	100 g.
No. 3			1500 g.
a ∫Stearin	90 g.	No. 6	
". Coconut Oil	10 g.	Glycol Stearate	30 g.
Caustic Potash (50° Bé.)	42 g.	Absorption Base (Parachol)	
b. Glycerin	20 g.	White Beeswax	30 g.
Water	100 g.	Sesame Oil	800 g.
c. Stearin	10 g.	Water	600 g.
As in No. 1.		l Saponine	16 g.

Shaving Creams, Non-	Foaming
Formula No. 1 (For Fa	itty Skin)
Stearin	50 g.
a. Stearin Vascline	10 g.
b. Triethanolamine Borax Water	15 g.
b. { Borax	1.5 g.
l Water	130 cc.
c. Alcohol (Perfume)	3 g.
Pour a, 70° C., into b,	60° C. Cool
stirring; add c before solids in collapsible tubes.	fication. Pack
No. 2	
Stearin	45 g. 2.5 g.
Triethanolamine	2.5 g.
Glycerin	15 g.
Water	67.5 cc.
Witch Hazel	50 cc.
Method as in No. 1.	

Latherless Shaving Cream U. S. Patent 1,991,501

A neutral shaving preparation of a latherless type which consists of a mixture of the following ingredients in substantially the proportion stated, steame acid 11 g., lanolin 10 g, coconut oil 0.3 g., concentrated ammonium hydroxide 1.35 g., paraffin wax 6 g., spermaceti wax 2 g., borne acid 15 g., water 75 g., and having a trace each of menthol, camphor and perfume.

Steame acid and hydrous landlin containing 20% water, together with coconut oil are melted together, and to this mixture is added the concentrated ammonium hydroxide, which contains approximately 25% of ammonia.

The waxes are then added and heating is continued until the entire mixture is hquefied. The resulting mixture is subsequently removed from the heat and a warm solution of the boric acid in approximately 75 g. of water is added while continuously stirring.

At this point, or at any point previously, the menthol, camphor and selected perfumes are added in amounts which give the most pleasing effect.

The mixture is then violently stirred until cold, and the final resulting product is a white cream.

Shaving Creams,	Nor	1-F)an	ing
Formula	No.	1		
Stearin				75 g.
a. Stearin Vaseline				13 g.
b. Triethanolamine Borax Water			٠	2 g.
b. { Borax		*		2 g.
l Water				195 g.
c. Alcohol				6 g.

Melt up a to 70° C., mix b and heat up to 60° C, then pour a into b with stirring. Shortly before the cooling (solidification) add perfume in the alcohol c, stir until cold. Fill into collapsible

2
36 g.
10 g.
5 g.
5 g.
130 g.
3
30 g.
10 g.
100 g.
45 g.
10 g.

Camphor Shaving Milk 50 g. Camphor, Spirits of 50 g. 2 g. Glycerin Lavender Oil 600 g. Alcohol Add: 25 g. Borax, Powder Distilled Water 1200 g. 200 g. Fresh Lemon Juice Stir; allow to stand over night; filter.

Milky-White Shaving Soap, L	iqui	d
Coconut Oil	30	g.
Tallow	90	g.
Stearie Acid	90	
Caustic Potash (50%) about	90	g.
Potassium Carbonate		g.
	370	
	120	
	210	
Perfume 2.5	-10	g.

Shaving Milks Formula No. 1

10 g.
2 g.
15 g.
40 g.
40 g.
10 g.
20 g.
20 g.
20 g.
440 g.
50 g.
40 g.
10 g.

92 1		AL FORMOLANT	
No. 3		Caustic Potash (50%)	bout 6.33 g.
Grind:			bout 6.33 g.
Lanolin, Pure, Pale	50 g.	Distilled Water (or	70
Coconut Oil	25 g.	Softened Water)	79 g.
Borax	8 g.		-
Neutral Soap Powder	25 g.	Shaving Soap, L	iauid
Water	80 g.	Olein, Light	9 g.
Rose Water	400 cc.	Coconut Oil, Cochin	3 g.
Orange Flower Water		Caustic Potash (50° Bé	
(Tepid)	400 ec.	Alcohol	1 g.
Peppermint Oil	2 ec.	Glycerin, C.P.	8 g.
		Water	73 g.
Astringent After Shar	ing Milk	Rose Water	1 g.
Formula No.	1	Shaving Soap, Similar to	"Rasibloc"
Glyceryl Monostearate	10 g.	(Stearin	100 g.
Vegetable Oils	8 g.	a. Glycerin	_
White Paraffin Oil, Odor	less 2 g.	1	- 6
Distilled Water	73 g.	b. Caustic Potash (39°	
Acetic Acid (50%)	5 g.	Caustic Soda (37° Bé.	.) 11.4 g.
Glycerin (28° Bé.)	2 g.	c. Coconut Soap	30 g.
Add perfume resistant t		Warm each portion and	l mix together
No. 2		in above order.	
Camphor	2 g.	After Shave Lo	tion
Eau de Cologne Oil	4 g.		
Alcohol	300 g.	Alcohol (95%)	680 g.
Glycerin (28° Bé.)	80 g.	Perfume Oil	6 g.
Rose Water	614 g.	Glycerin	15 g.
	-	Tannie Acid	5 g.
Transparent Liquid Sh	aving Soap	Distilled Water To the alcohol perfum	294 g.
Olein, Clear, Pale		glycerin, then the water-ta	
Coconut Oil	1.575 g.	tion.	mile acid soid

POWDERED HAND TOILET SOAPS

No. 1

Formula:

Dry Yellow Powdered Soap, 92% plus c.p.s.,* S.N.† to be over 210 titre,‡ 25 to 35° C
Cocoanut soap-powder, 30% Anhydrous
Soap Contents, S.N. to be over 210
titre, 30 to 35° C
Wyo-Jel No. 719 (Colloidal Bentonite),
200 mesh
Tri-Sodium Phosphate, tech. grade pow-
dered
Perfume
Citrene
Girella
Camphory Sassafras Oil

- c.p.s. = Chemically Pure Soap. † S.N. = Saponification Number. † Titre = Melting Point of Fats.

Bathroom Factory Office and Traveland and Ga-Dispenser Home Use rage Use General 75 lb. — 40 lb. 60 lb. --- 60 lb. 25 lb. 20 lb. 24 lb. 33 lb. 30 lb. 20 lb. 1 lb. 7 lb. 5 lb. 0.2 lb. 0.1 lb. 0.1 lb.

0.7 lb.

No. 2

No. 3

No. 4

The ingredients are weighed into a clean and dry mixer and intensely mixed for 15 to 20 minutes. The perfume should be sprayed or sprinkled over the powdered soap or soap-powder to avoid caking. As none of the ingredients are hygroscopic it is not necessary to pack the finished product air tight.

For starting production, a clean openhead steel drum rolled and shaken on the floor is satisfactory for mixing, providing some wooden weights are laid inside to assure agitation. However, for big scale production, use one big horizontal mixer, 2000 lb. capacity, cylinder driven from both end countershafts and equipped with a double action agitator which moves toward the 6" x 8" outlet in the middle and which is driven by a 15 h.p. motor. A slip ring motor, or a compensator allows this mixer to be started with a full load, thus avoiding accidents and dusting.

The most ideal process to make powdered hand toilet soaps is by making them wet-processed, and if other soaps are also manufactured, it is easy and much more preferable to do so. In the case of Formula 1, the Wyo-Jel is crutched into the hot molten soap stock before cooling and drying and the perfume is added immediately before grinding down of the dried soap flakes. In case of Nos. 2 and 3, paste soap, regular soap powder is hot mixed with all the ingredients added at once to a bakery-type dough mixer. In case of hot processing much more Wyo-Jel can be used and the final structure will be more uniform and much harder to duplicate.

Liquid Soaps (French) Formula No. 1

Olive Oil Soap:

(Caustic Potash (Solid) Water minimum possible for solution

182 kg. b. Olive Oil 362 kg. Palm Oil 362 kg. Coconut Oil Heat to 49° C., add to a.

c. Alcohol 170 l. Boil the whole under reflux (82° C.).

When saponified, cool, and add

5.6 l. d. Water

No. 2

Coconut Oil Sonp

a. Soda Ash 1 kg. Water 10 1. b. Wood Ashes 15 kg. Water

Extract through a tin can with holes, pouring through water 3 to 5 times.

c. Canstie Soda 1. Boil 10 to 15 min.:

I part by volume a. 4 parts by volume h. 6 parts by volume Add Coconut Oil 10 parts by volume

during the boiling in small parts, stir slowly. Then dominish heat, stir continuously, take off, stir, then pour into wooden forms.

2. Or: Boil 10-15 minutes:

4 parts by volume 6 parts by volume

Sodium Sulphate

(10%)1 part by volume Salt 1/2 part by volume Add: 9 parts by volume

Coconut Oil and after:

Tallow 1 part by volume

Method as in No. 1. Gentle boiling, thorough sturing, dry.

No. 3

Liquid Coconut Oil Soap

(Water 1. a. |Canstic Potash (Solid)

Add a to

b. Coconut Oil (49° C.) 20 kg. c. Alcohol 2.5 1.

Warm the whole to 82° C. under reflux as in 1. Let cool 24 hours, then add:

d. Water very little Potassum Chloride optional Glycerin

No. 4

Liquid Glycerin Soap

Soft Soap, Good 21 g. 7 g. **Glycerin** Water 14 g. Alcohol Tale or Pumice 5 g.

Let stand for several days; take care to eliminate excessive alkali by adding oleic acid. Filter.

Transparent Glycerin Soaps Formula

No.	1	No.	2	No.	3

Coconut Oil,			1 .
Cochin	20	26	30 kg.
Tallow	18	21	20 kg.
Castor Oil	12	10	15 kg.
Caustic Potash,		1	
40° Bé.	25		— kg.
36° Bé.		32	kg.
39° Bé.	_		35 kg.
Glycerin	10	13	10 kg.
Sugar	10	40	42 kg.
Water (60° C.)	15	30	38 kg.
"Fillers"	_	30	35 kg.

To this soap-base add distilled water in small portions to about 15 (kg.), and to the resulting clear, but very soft, soap add a hardening solution (of 15° Bé.), made up of:

Potassium Carbonate Sal Soda	1 kg. 1 kg.
Salt	1 kg.
Add water to get 15° Bé.	Warm to
,	

Add enough to get samples of suffi-ciently hard soap. Let stand covered for an hour, and test result.

Should not be of too high viscosity when spread on a glass-sheet. If too viscous or too foamy add water.

Add perfume at 50° C., sift in dye,

stir and pour into molds.

Transparent Soap (Without Glycerin)

Tallow, Cochin	24	
Coconut Oil	24	kg.
Castor Oil	16	kg.
Heat to 50-60° C.		
Add in thin jet:		
Caustic Soda (39° Bé.)	33	kg.
Stir until soap swims on cover. Stir slowly over water	top, bath.	thei Ado
Alcohol	1-2	kg.
than		

Water (60° C.) 22 kg. 20 kg. Sugar Again Alcohol 18-19 kg.

Alcohol

Cover. Keep at 75° C. for an hour.

Soap should be dark and clear; foam light. Soap should remain "knife-thick" on a glass-sheet.

If opaque, try (before in test-tube) to add slowly hot water, or caustic soda (20° Bé.).

At 50-60° C. add perfume and the last 3-4 kg. of phone alcohol

3-4 kg. of above alcohol.

Rose Soap

	0,000 g.
Cinnabar, Moistened 6	60-80 g.
b. Rose Essence	25 g.
Geranium Essence	60 g.
Clove Essence	15 g.
Chinese Cinnamon Essence	10 g.

Palm Soan

	r ann soap			
a.	Pure Palm Soap	5000		
	Half Palm Soap	5000	g.	
b.	Bergamot Essence	60	g.	
	Chinese Cinnamon Essence	25	g.	
	Clove Essence	15	g.	
	Essence of Fine Lavender	30	g.	

Althaea (Marshmallow) Soap

a.	White Tallow Soap	5000	g.
	Pure Palm Soap	5000	g.
b.	Yellow Ochre	30	
	Paris Red	30	g.
c.	Essence of Fine Lavender Essence of Pressed	15	g.
	Lemon Peel	16	g.
	Essence of Neroli		

16 g. Petitgrain Essence of Verbena 10 g. Essence of English Mint

Bouquet Soap

a.		10,000	
	Brown Ochre	100	g.
b.	Essence of Bergamot	80	g.
	Essence of Cloves	15	ġ.
	Essence of Neroli	15	g.
	Essence of Sassafras	10	g.
	Essence of Thyme	10	g.
or	also:		
b.	Essence of Fine Lavende	r 20	g.
	Essence of English Mint	20	g.
	Essence of Pressed		
	Lemon Peel	25	g.
	Essence of Sage	20	g.
	Essence of Thyme	10	g.

The following Soaps using Lauryl Sulphonates are covered by German Patents.

I. True Lemon Soap

Citric Acid	5 g.
Sodium Citrate	1 g.
Lanolin-Vaseline Oil (2:1,	_
1:1)	5 g.
Vegetable Lecithin	2 g.
Glycerin	2 g.
Lauryl Sulphonate	85 g.
Lasury i Burphonate	OU 5.

		COMETICS	AND DRUGS 95
	Liquid Tar Soa	n	V. Chlorthymal Sonn
Wood Tar (9 3 g.	V. Chlorthymol Sonp Chlorothymol 1 g.
Glycerin	20,00	5 g.	Chlorothymol 1 g. Acetic Acid, Concentrated 0.5 g.
Triethanolan			Alcohol 3.5 g.
Sulphonat	e	92 g.	Triethanolamine Lauryl Sulphomite 95 g.
II	I. Alum Soap		
Aluminum S		5 g.	VI. Chlorine Soap Chloramm 1-2 g.
Lorol Sulpha	te or Tri-	* B*	Landm Paraffin Od (1:1) 5 g.
	ne Lauryl Sul-	0"	Glycerm 3 g.
phonate		95 g.	Sodium Lauryl Sulphonate 90 g.
IV	. Iodine Soap		And the same of th
Iodine-Alcoh	ol Solution	5 g.	Soap for Removing Scarred Skin
Glycerin	ing Laurul	10 g.	Liquid Paraffin 70 cc. Medicated Soap, Powdered 70 g.
Triethanolan Sulphonate		85 g.	Sodium Peroxide 21/2 up to 10 g.
~ u-r		POWDER I	
		Mag	
	Rico	neson Colloidai Car-	i Mag- nesium Zine Cold Other
	Starch Talcum		Stearate Oxide Cream Additions
Face Powder:	600 200	. 100	40 60
	450 300 500 300	50 25	220 70 Titanium Dioxide
	500 300	100 250	5
Body Powder:	900		. 90 10 Salicylic Acid
	70 850	70	20 10 100 Boric Acid 60 Boric Acid
	80 490	300	100
Infant Powder	1000		. 6 1 Lanolin
Foot Powder:			
	850		100 10 Salicylic Acid
	800		200 100 Boric Acid
	750		260 350 Kieselguhr 10 Thymol
	600		10 Thymol 0 1 Formaldehyda
Di	isting Powders		No. 6
ŀ	ormula No. 1	_	Bismuth Subgallate 5 g. Boric Acid 15 g.
Phenol		1 g. 3 g.	Borne Acid 15 g. No. 7
Camphor Exsiccated .	Alum	96 g.	Bismith Submitrate 20 g.
man attent.		.,	Starch 10 g.
~ " " 1	No. 2	4 g.	Purified Tale 70 g.
Salicylic Ac Baric Acid	10	5 g.	No. 8
Starch		16 g.	Mercuric Chloride 0 06 g. Sodom Salicylate 26 g.
Purified Ta		60 g.	Sodium Salicylate 26 g. Prepared Chalk 4 g.
	No. 3	10	
Salicylic Ac Bismuth Su	id haiteata	10 g. 15 g.	Throsulphate Dusting Powder
Zinc Steara		10 g.	Sodium Throsulphate 6 g.
	No. 4		Boric Acid 24 g.
Salicylic Ac		2 g.	Dusting powder (prophylactic) for
Tannoform		13 g.	ringworm.
Talcum	No. 5	15 g.	77 . 77 . 17 . 17
Salicylic A		2 g.	Foaming Bath Powder
Tannic Acid		5 g.	Sodium Acid Carbonate 40 g. Starch, Wheat 50 g.
Orris Root		33 g. 60 g.	Sodium Carbonate 10 g.
Mulli		0	*

Tartarie Acid	30 g.	Clove Oil	5 cc.
Kaolin, Colloidal	20 g.	Fennel Oil	5 cc.
Soap Powder, Concentrat		Ceylon Cinnamon Oil	1 cc.
		Lemon Oil	
Saponin	5 g.	Lemon On	1 cc.
Keep completely dry and	l scaled from		•
air to avoid decomposition	. 1-2% per-	Oxygen Tooth P	asta
fume (Lavender, Pine Ne	edle. Éan de		
Cologne, Fancy), is added.		Calcium Carbonate, Prec	
cologic, ranej,, is added.		tated, Medium Density	
	1	Glycerin, C.P.	30 g.
Mentholated Talc	um	Hard Fat Soap Powder	7 g.
Menthol			il soft paste
	0.25 g.		•
Alcohol	5 cc.	To 100 parts of this par	•
Talcum	50 g.	Sodium Perborate	10-15 g.
Dust freely on itching pa	rt. 1	Perfume	1 g.
	_		
"Prickly Heat" P	owder	Tale Tooth Pa	ste
Starch	121/2 lb.	Purified Talc	42 lb.
Tale	7 lb.	Magnesium Carbonate	8 lb.
Zinc Stearate	1/2 lb.	Phenol	1/2 lb.
		Two on on mth	
Camphor	2 oz. 5 lb.	Phenol Tragacanth	
Zine Oxido		On or Ominge	2½ dram
Menthol	1 oz.	Oil of Lemon	5 oz.
		Oil of Anise	1 dram
Tooth Paste		Oil of Peppermint	6 oz.
		Menthol	5 oz.
Soap Powder	2500 g.	Glycerin	6 gal.
Calcium Carbonate	500 g.		-
Lactose	150 g.		
Glycerin (28° Bé.)	2000 g.	Salt Tooth Pa	ste
Water	400 g.		
Peppermint Oil	100 g.	U. S. Patent 1,90	18,888
Alcohol	100 g.	Glycerin, C.P.	37½ lb.
	100 g.	Neutral Soap	11/2 lb.
Carmine	10-20 g.	Gun Tragacanth	1½ lb.
		Magnesium Carbonate	172 10.
Tooth Paste with Low Gly	cerin Content	(Einstein Carbonato	13 lb.
Calcium Carbonate, Prec		(Finely Divided)	19 10.
tated, Medium Density	45.0	Calcium Carbonate	F11/ 11
		(Finely Divided)	51½ lb.
White Clay (Bolus Alba		Milk of Magnesia (Mag	; -
Soap Powder (85-88%),		nesium Hydroxide)	31 lb.
Pale, no Odor or Taste	10 g.	Distilled Water	24 pt.
Water	20 g.	Saccharin Powder	282 gr.
Glycerin, C.P.	20 g.	Salt (Finely Divided)	108 Ìb.
Month Donto (With	Clusterin)	Flavor	
Tooth Paste (Without		Menthol Crystals	2% oz.
White Clay (Bolus A	dba) 30 g.	Oil of Peppermint, U.S.	P. 8 0z.
Calcium Carbonate	-	On of reppermint, U.S.	2/ 05
a. Precipitated	15 g.	Oil of Anise, U.S.P.	2/₃ oz.
Soap Powder (as Abo		Methyl Salicylate	$\frac{2}{3}$ oz.
b. Tragacanth Paste (19	%) until pastv	*Flavor Compound	
Baccara - abov (1)	· / 2 F	No. 04595	12 oz.
Mark Street Stre	-	* Flavor Compound No. 04	595 is comprised
Tooth Paste		as follows:	-
Calcium Carbonate, Pre-		Twice Rectified Oil of	
cipitated	50 g.	Peppermint	274 02.
White Bolus	10 g.	Oil of Eucalyptol Oil of Wintergreen	90 oz. 45 oz.
Glycerin (sp. gr. 1.24, 30	° Ba) 20 g	Rectified Aniseed Oil	22 1/4 02.
Water	18 g.	Safrol	22 1/2 02. 22 1/2 02.
		The glycerin, water, so	
Tragacanth	1 g.	canth, milk of magnesia,	and saccharin
Perfume (as below)	F0	cantil, milk of magnesia,	ivor
Peppermint Oil	50 cc.	are mixed with a rapid n	HACE.
Menthol	5 cc.	Then flavor is added, w	
Anise Oil	25 cc.	I made a few days in adva	nce, and arter

15 minutes of mixing the product is
transferred to a small mixer, the salt is
added, the mixer is run for five minutes
more, then the magnesium carbonate is
added, followed by another five minutes'
run, after which the calcium carbonate is
fed to the pasty mass, and, after this
has been taken up, the batch is run for
20 minutes more.
no institute and the

The finished mass is allowed to stand for 12 hours, and, after stirring slowly for 10 minutes before filling, the mass is filled into ordinary collapsible tubes.

Denture (Artificial Teeth)	Clear	er
Glycerite of Starch	36	g.
Diglycol Laurate	1	g.
Sugar Syrup	2.25	
Magnesium Carbonate	1.13	
Gum Tragacanth	.07	g.
Precipitated Chalk	41	g.
Sodium Bicarbonate	6	g.
Water	10.5	g.
Flavor	to s	uıt
Denture (Artificial Teeth)	Adhere	nt
Gum Karaya	80	ø.
Gum Arabic	20	
Cum Arabic	_	0
Dental Impression Mat	crial	
British Patent 399,8	42	
Copal	26	g.
Stearic Acid	21	g.
Shellac	5	g.
Melt together and then add	while	heat
ing and stirring:		
Talc	48	g.
Iron Oxide, Red		g.
Iron Oxide, ited	/1	
Temporary Dental Fil	ling	
Zinc Oxide	85	g.
Rosin, Powdered	15	g.
Oil of Cloves		g.
Canada Balsam	35	ø.

Dental Canal Cement

35 g. 5 g.

Thymol	1 μ	z
Rosin	9 8	ż
Chloroform	150	
CHIOIOIOIM	_	-

Canada Balsam Peru Balsam

Dental Pulp Capping

Make a paste of zinc oxide and eugenol.

Dental Pulp Devitalizer

Make a paste of arsenic trioxide and
eugenol

Antiseptic Mouth Wash ("Listerine" Type)

11.1	
50	g.
1	g.
1	g.
0.125	cc.
0.5	cc.
	cc.
0.1	cc.
250	cc.
1000	ee.
to co	olor
	50 1 1 0.125 0.5 0.25 0.1

The boric acid is dissolved in the water or about 700 ec. of same. All the other products are dissolved in the alcohol and the two solutions mixed and colored to a very pale straw. The above product must be labeled 25% grain alcohol.

Mouth Wash Tablets

2.20.000	
Peppermint Oil	30 cc.
Saponin, Best	100 g.
Sodium Benzoate	500 g.
***************************************	_

Mouth Rinse

Salt	30 g.
Sugar	20 g.
Oil of Cinnamon	1/4 ec.
Oil of Cloves	⅓ cc,
Oil of Peppermint	¼ cc.

Gingivitia Month Wash

tingities acousti	*** 64.07**
Boric Acid	4 g.
Potassium Chlorate	8 g.
Peppermint Water	350 cc.

Breath Deodorant

Dissolve one 4.6 grain tablet chloramine in 1 oz. water. Brush teeth and tongue, and rinse out mouth with this solution, while fresh.

Immediately and permanently rids breath of even such odors as those of garlic and onions.

Depilatory

German Patent 601,078

Barium Sulphide	100 oz.
Starch	60 oz.
Magnesium Silicate	30 oz.
Pyrogallol	10 oz.
Make into a paste with	water before
using.	

Odorless Depilatory

Pernyaroi	0.0-0	ъ.
Polychol (or Polyglycol)	5	
Lanolin Anliydrous	20	g.
Rub together till uniform.		

Adhesive Depilatory U. S. Patent 2,013,928	Sunburn-Protecting Oil Quinine Oleate, C.P. 3-5 g.
Rosin 90 g.	Paraffin Oil 27 cc.
Cottonseed Oil 10 g.	Fatty Oil 70-68 cc.
Warm together and stir until uniform.	Dye (Oil-Soluble Red)
	Married Confession of the Conf
Sun Burn-Protectors	Sunburn-Protecting Oil
Liquid	Quinine Ricinoleate 3-5 g.
a. Triethanolamine 40 g.	Olive Oil 97-95 cc.
Trihydroxyethylamine	to the second se
Stearate 40 g.	Sunburn (Suntan) Oil
Melt on water bath, make emulsion in	'Mix
Water (60° C.) 620-630 g.	Vaseline Oil 75 g.
b. Paraffin Oil 100 g.	Sesame or Peanut Oil, Pale 23 g.
Peanut Oil 150 g.	Thymol 0.5 g.
Oleic Acid 30 g.	Lanolin, Anhydrous 1.5 g.
Warm up on water bath to 40° C. Methyl p Hydroxy Ben-	Perfume 1-2 g. Made up of:
zoate 1 g.	Pine Oil 3 cc.
Pour b into a, perfume with	Lavender Oil 1 cc.
c. Perfume Oil to suit	Rosemary Oil 1 cc. Laurel Oil 3-5 cc.
*Stir until cold.	
Cream	Suntan Oil
White Wax 60 g.	Paraffin Oil 20 cc.
Cocoa Butter 30 g.	Fatty Oils, Free from Acid,
Lanolin, Auhydrous 40 g.	Preserved 80 cc.
Peanut Oil 300 g.	Etheric Oils (Bergamot, Eau de Cologne [free from
Spermaceti 20 g.	Methylanthranilic Ester] or
Moldex or Other Preservative 1 g. Perfume 5-10 g.	Pine Needle Oil) 1 cc.
	Dye with Chlorophyll, Oil-soluble.
Preventatives against Sunburn	
a. Gum Tragacanth (Powder) 15 g.	Preparations to Protect Feet Against
Glycerin 50 g.	Hurting and Inflammation
Grind in mortar.	Foot Creams
b. Quinine Acid Sulphate 100 g. Citric Acid 100 g.	Formula No. 1
Citrie Acid 100 g.	Potash Soap 50 g.
Water 1200 g.	Yellow Vaseline 15 g.
Alcohol (95%) with Perfume 400 g.	Water 29 g. Zine Oxide 6 g.
c. Glycerin 150 g.	Zine Oxide 6 g. Caustic Soda 11 drops
Grind a, then add the b solution, and	No. 2
finally add c.	Potash Soan 52 o
	Vaseline 15 g.
Sunburn Protecting Cream	water 21 g.
a. Quinine Hydrochloride 4 g.	Zinc Oxide 6 g.
Ålcohol (95%) 12 g.	No. 3
b. Citric Acid 0.8 g.	Soap 35 g. Vaseline 15 g.
Water 10 g.	Vaseline 15 g. Water 45 g.
c. Tragacanth Powder 3.5 g.	Zinc Oxide 5 g.
Glycerin 10 g.	Lavender Oil to suit
Water 42.5 g.	No. 4
Mix solutions a and b and then work into solution a	Lamb Tallow 100 g.
Perfume Composition, with	Pig Fat 100 g. Creosote 1 g.
Fresh Perfume Odor 9 drops	Juniper Oil 10 g.

No. 5	
Wool Fat	20 g.
Vaseline	10 g.
Formalin	10 g.
No. 6	
Glyceryl Monostearate	20 g.
Glycerin	5 g.
Paraffin Oil	5 g.
Formaldehyde Solution	15 cc.
Water	55 cc.
Melt up to 60° C. Stir until	l cold.

Pecling Paste for Corns or Hard Skin (Not to be put on normal skin, as it is irritating).

Formula No. 1

Lard		50 g.	
Salicylic Acid, U.S.P.		50 g.	
No. 2		•	
Salicylic Acid, C.P.		30 g.	
Vascline, White		70 g.	
No. 3			
Mild-acting paste (stir	war	n):	
a. Pine Resin, Pure Wax, Yellow Larch Turpentine Vaseline, Yellow	8	g. } Mel	
Wax, Yellow	30	g. S Mei	ı
a. Larch Turpentine	12	g.	
l Vascline, Ŷellow	16	g.	
b. Salicylic Acid	8	g.	
Anaesthesin	3	g.	
Ponnut Oil	14.5	σ.	

b. Salicylic Acid 8 g.
Annesthesin 3 g.
Pennut Oil 14.5 g.
Mix warm, stir until clear solution; cool stirring; when thickening starts, add
Methyl Salicylate 0.5 g.
Peru Balsam 8 g.
Stir until cold.

Athlete's Foot Ointment

Salicylic Acid	8 oz.
Ammoniated Mercury	4 oz.
Bismuth Subnitrate	12 oz.
Oil of Eucalyptus	12 oz.
Hydrous Wool Fat	64 oz.
Mix and make into an	ointment.

Athlete's Foot Powder

Sodium Thiosulphate	20 07.
Boric Acid	50 oz.
Purified Talc (Sterilized)	30 oz.
Triturate thoroughly. This	may be
used as a prophylactic powder a	pphed to
the feet and dusted in the shoes	

Athlete's Foot Treatment

Immerse feet two or three times a day in a warm saturated aqueous solution of furfural. Always have a little free furfural floating around to make sure of

an excess. Continue treatment until all signs of the disense disappear. Then treat feet once a day for several weeks to prevent recurrence. Shoes and socks should also be treated with this solution to disinfect them.

"'Athlete's Foot" Remedy

Gentian Violet	1 part
Alcohol	100 parts
Water	100 parts
12.2 (1.12. 1.1)	

Stir until dissolved.

	Bunion	Remover	
Salicylic	Acid		6 g.
Lanolin			60-g.
Quality for	d in hot	mater 411	off thick

Soak foot in hot water; cut off thick skin and apply twice a day.

Pilocarpine Eye Drops

Pilocarpine Nitrate	0.1 g.
Borne Acid	0.2 g.
Distilled Water	to make 10 cc.
Label: Drop into eye	from one to five
imes daily (in chronic	ghucoma).

Pilocarpine Eve Salve

Pilocarpine Nitrate	0.2 g.
Distilled Water	1 cc.
Hydrous Wool Fat	2 g.
White Petrolatum	7 g.

Mix with careful trituration and dispense in collapsible tube with eye tip.

Label: Apply to affected eye at hedtime (in chrone glaucoum). If collapsible eye outment tube is not available, a glass rod may be used to apply salve to lower lid, which is then permitted to close. Gentle massage of lids helps to distribute ointment over the conjunctiva-

Eve Ointment

,	
Silver Nitrate	0.5 g.
Distilled Water	1 g.
Cocoa Butter	15 g.
Liquid Paraffin)	equal parts
Soft Paraffin	to 100 g.

Cetyl Alcohol

U. S. Patent 2,021,926

Formula No. 1

241 parts of spermaceti are melted and heated to 200° C. 42 parts of powdered potassum hydroxide are now added with agitation in half an hour, during which time the temperature is allowed to rise to 240° C. It is held at this temperature for half an hour when superheated steam

is passed in. There distils over with the steam a colorless oil which sets on cooling to a crystalline waxy solid which is entirely free from fatty acid and from unsaponified spermaceti. The yield is approximately 100 parts by weight, the proportion of water to oil in the distillate being approximately 10:1.

241 parts of spermaceti are treated as in Example 1 with a mixture of 21 parts in Example I with a mixture of 21 parts powdered potassium hydroxide and 15 parts of powdered sodium hydroxide. After reaction, the molten mixture of soaps and fatty alcohol is subjected to superheated steam distillation at about 250° C., eventually at 280° C. until no more oil distils. The yield is approximately 100 parts of the pure alcohol from spermaceti, the ratio of water to oil in the distillate being approximately 10:1.

No. 3

268 parts of sperm oil are treated as in Example 1 with a mixture of 21 parts of caustic potash and 15 parts of caustic soda. After reaction the mass is sub-jected to superheated steam distillation until no more oil distils. The yield is 90 parts of a semi-solid alcohol, free from unsaponified wax or free fatty acid. The ratio of water to oil in the distillate is approximately 4:1.

Arthritis Ointment

20 g. Ichthvol 30 g. Lanolin

Rub together until uniform; apply freely to joint and apply bandage.

Frostbite Ointment

Ichthyol		3	g.
Ichthyol Lanolin			g.
Camphor		2	g.
Petrolatum		60	g.
		dissolved.	Ruk
i-41-1 1	1		

into skin and bandage

Analgesic Balm

Menthol	5	oz.
Methyl Salicylate	10	oz.
Hydrous Wool Fat	75	oz.
White Petrolatum	10	oz.

Burn Ointment

Tannic Acid	2	g.
Ichthyol	33	
Lanolin	62	g.

Carbuncle Ointment

Ichthyol Lanolin	25 g. 35 g.
Zinc Oxide Ointment	90 g.
Apply thickly daily.	

Chapped Skin Ointment

Phenyl Salicylat	e		8	g.
Menthol			4	
Olive Oil			40	
Lanolin			125	g.
Warm together	and	mix	until	dia
solved.				

Glycerin-Sulphur-F	Caolin-Acne Paste
Kaolin	10 g.
Sulphur, Colloidal	7.5 g.
Glycerin (21%)	to pasty consistency

Boil Ointment

Ichthyol Lanolin			15 g	
Apply thickly on	gauze	and		
place with adhesive.				

Ringworm Ointments

Sulphur Ointmen	.t	
Precipitated Sulphur	1.	5 g.
Petrolatum	30	g.
Rub in gently once or	twice	daily.

Strength may gradually be increased up to 20 per cent.

Compound Benzoic Acid Oin	tment
Salicylic Acid	1 g.
Benzoic Acid	2 g.
Ointment of Rose Water	30 g.
	Strength
nay be doubled, if necessary.	
Chrysarobin Ointment	

Chrysarobin 1.5 g. Petrolatum Apply with care against getting it in the eyes.

Salicylic Acid Pigment O-1:--1:- A .: 3

Sancyne Acid	1.0	к.
Chloroform	30	cc.
Paint on affected area twice	daily	until
descus mation occurs		

ryrethrum Omtment	
Pyrethrum Extract	27 g.
Absorption Base (Parachol)	73 g.
Mix until smooth. Useful in	treating
scabies and other insect infesta	tions.

Ulcer Salve	
Ethyl Aminobenzoate	3 g.
Paraffin	10 g.
To to latum	20 σ

Petrolatum Spread on gauze and apply to ulcer.

Protecting Skin Against Mustard Gas Glycerin impregnated coarse fibered clothing is recommended. This protection lasts for at least two hours' exposure to this gas.

A. B. C. Liniment

Tincture of Aconite	30 cc.
Fluidextract of Bell	
Chloroform	30 сс.
Soap Liniment	to make 240 cc.
Analgesic liniment.	For external use
only.	

Glycerin-Sulphur Limment

20 g.
20 g.
20 g.
20 g.
20 g.

Penetrating	Dilling	
Oil of Turpentine	1	gal.
Oil of Sassafras	1	lb.
Oil of Cajaput	1,,	1b.
Chloroform		gal.
Oil of Camphor	5	97.
Oleoresin Capsicum		gal.
Coal Oil		8

Rheumatism Liniment 1 lb.

Campnor		
Chloroform	32	fl. 07
Alcohol	80	fl. oz.
Methyl Salicylate	16	fl. oz.
Dissolve camphar in th	e mixtur	e of th

other ingredients. Excellent for sore or aching muscles. Should be applied at night by rubbing in.

Back Rub Ointment

Zinc Stearate	5	g.
Tincture of Benzoin	5	g.
Scarlet Red Omtment	(5%) 0.25	g.
Hydrous Wool Fat	.50	g.
Liniment of Camphor	180	cc.
Mutton Tallow	500	g.

Non Staining (Non Leaking) Mineral Oil Laxative

White Soft Paraffin Wax	2 oz.
Mineral Oil, U.S.P.	6 oz.
Warm together and stir until	uniform

Castor Oil Candy Laxative U. S. Patent 1,991,139

Predetermined quantities of broken chocolate and castor oil are heated in separate containers or kettles before mixing. The chocolate is heated to approximately 115° F., while being thoroughly stirred or agitated, and is then permitted to cool to approximately 85° F., which temperature is finally slowly increased to between 88 and 90° F.

After the chocolate melting operation has been commenced, or simultaneously with this operation, an amount of castor oil approximately that of the melted on approximately that of the latter of the chocolate, is slowly heated to between 85 and 90° F., preferably between 88 and 90° F. The heating of the castor oil and chocolate is so timed that the temperature of the one will coincide with that of the other. The best mixing tempera-ture is between 88 and 90° F., it being essential that the temperature of each ingredient be kept exactly the same.

Mixing of the melted chocolate and heated castor oil is effected at this stage by drawing off the two ingredients from their separate kettles into a mixer, where they are thoroughly beaten and blended, after which the temperature is lowered to between 75 and 80° F. At this point, the mixture is cust into centers or chocolate shells which are subsequently capped with chocolate and run into a cold box for final cooling.

Agar Mineral Oil Emulsion

gai.
lb. lb. oz.

Some sodium benzoate and aseptoform as preservative, and a small amount of vanillin and saccharin for flavoring pur-

Emulsion of Liquid Petrolatum Liquid Petrolatum 500 Acacia, in Very Fine 125 Powder 100 Syrup Vanillin 0.035 g. 60 Alcohol Distilled Water, a sufficient quantity to make

Mix the liquid petrolutum with the powdered acacia in a dry mortar, add 250 cc. of distilled water all at once and emulsify the mixture. Then add, in divided portions and triturating after each addition, a mixture of the syrup, 50 cc. of distilled water and the vanillin, dissolved in the alcohol. Finally add sufficient distilled water to make the product measure 1000 cc.

Note: In preparing Emulsion of Liquid Petrolatum other methods of emulsification may be used and the quantity of acacia may be reduced or it may be replaced by agar, gelatin, tragacanth or mixtures of any of these emulsifying agents, provided the resulting emulsion is similar in viscosity and appearance to the emulsion made by the formula suggested above.

Antipyrine Suppositories

Antipyrine	3 g.
Extract of Belladonna	0.1 cc.
Cacao Butter	20 g.
Mix and divide into ten	suppositories

Mix and divide into ten suppositories.

Label: One every two to four hours as required.

Psoriasis Treatment Formula No. 1

Salicylic Acid 10 g.
Oil of Cade 25 cc.
Soft Soap 25 g.
Alcohol to make 100 cc.

Paint over patches, permit to dry, and wash off excess in bath.

No. 2

Salicylic Acid	10 g.
Chrysarobin	20 čc.
Oil of Cade	20 cc.
Soft Sonp	25 g.
Petrolatum	25 g.
Labels Apply to notaboo	

Label: Apply to patches.

Acidosis Preventative

To a teaspoonful of sodium bicarbonate in a deep bowl, add the juice from one lemon. Stir until effervescence is completed, and add a glass of cold water, and driuk. Best results are obtained by taking this drink upon rising in the morning, at least one-half hour before breakfast.

Cold and Grippe "Remedy"

The following has been used with splendid success by members of a technical manufacturing organization:

a. Acetic Acid (36%),

U.S.P. ¼ fl. oz. Water to make 1 fl. oz. b. Ammonium Carbonate, U.S.P. 48 gr. Water to make 1 fl. oz.

c. Sodium Bicarbonate 2 d.
Potassium Citrate 2 d.
Aromatic Spirits of
Ammonia 1 fl. oz.
Water 1 fl. oz.
Mix a and b; after effervescence stops add c.
Take one teaspoonful every 2 hours.

Hay Fever Remedies

roimina No. 1	
Ephedrine (Dried)	0.1
Petrolatum, Liquid	10 c
Use as nasal spray.	

No. 2 Ephedrine Sulphate

Calcium Lactate 4 g.
Place in No. XXX capsules; use one
3 or 4 times daily.

Sea-Sickness Remedy

Antipyrin Sodium Bromide Sugar		4 g. 8 g.
ougai		2 g.

Use once every three hours.

Appetite Stimulant

Tincture of Capsicum	2	cc.
Tincture of Nux Vomica		cc.
Tincture of Gentian Compound	72	cc.

Doso: Three teaspoonfuls daily.

Bronchitis Inhalant

Menthol	1/2	g.
Chloroform	4	cc.
Tincture of Benzoin	120	cc.
Inhale twice daily, using	one teas	poon.

ful to pint of boiling water.

Menthol Inhalator

Eucalyptus Oil	4	cc.
Menthol	2	g.
Paraffin Oil	94	ec.

Laryngitis Spray

Thymol	0.15 g.
Menthol	1.2 g.
Eucalyptus Oil	3 g.
Petrolatum, Liquid	300 cc.

Tonsilitis Gargle

- Olibratio Guigio	
Potassium Chlorate	8 g.
Tincture Ferric Chloride	12 cc.
Glycerin	60 cc.
Water	240 cc.

	COSMETICS
Stomach Gas Re	lie f
Calomel	3 g.
Bicarbonate of Soda	3 g. 5 g.
Lactose	4 g.
Periodic Pain Alle	viator
Formula No.	1
Amidopyrine	20 oz.
Alcohol	40 oz.
Simple Syrup	138 oz.
Flavor	to suit
No. 2	
Starch	90 oz.
Amidopyrine	90 oz.
Acetyl Salicylic Acid	25 oz.
Camphor Tablets (Phari	naceutical)
Camphor	5 g.
Sugar	50 g.
Peppermint Oil	2-2.5 cc.
Pack tight, to prevent v	olatilizing.
Moth Protection T	'ablets
	225 g.
Naphthalene	75 g.
Camphor	50 g.
Ceresin	
Melt together and then a	aa
11 11	50 a

50 g. Hexachlorethano 5 g. Pine Needle Oil Dip cardboards into the above while

Sterilizing Helmets and Gas Masks

The U. S. Government, in its specifications for sand blast helmets purchased by its various departments, requires that each article be capable of passing either one of the following sterilization tests:

- (a) Immersion for ten minutes in a solution of formaldehyde made by placing one part of 40% solution of formaldehyde in nine parts of water, or
- (b) Subjection to sterilization by a moist atmosphere of antiseptic gas, preterably formaldehyde, for a period of ten minutes, at room temperature.

It has been suggested that care should be taken to remove all the formaldehyde from the masks by washing with water before they are placed in use.

"Creolin" Disinfectant

100 kg. Sulphonated Castor Oil Caustic Soda (36° Bé.) 51.2 kg Heat above at 80-100°C., then add 51.2 kg. 104 kg. Rosin Mix with heating until uniform and Tar Oils (200-320° C.) 775 kg.

Mix until dissolved and then add Water to make 1000 kg.

Disinguation for Malout

Disinfectant for Telephones			
Solution 1			
Oil of Wintergreen	0.5	g.	
Oil of Eucalyptus	0.5 0.25	g.	
Denatured Alcohol	15	g.	
Solution 2			
Formaldehyde	25	cc.	
Water	225	ce,	
Add solution 1 to solution	2 and	dilute	

Counter Irritant, Extra Strong

with water to 1000 cc.

Menthol				2 g.
Volatile	Oil	of	Mustard	2 cc.
Alcohol				50 cc.

Apply a few drops to affected area. (Must not be used in the vicinity of the eyes.)

Stainless Iodine Solution

Resublined Iodine	4 g.
Potassium Iodide	10 g.
Hyposulplute of Soda	10 g.
Alcohol, Anhydrous	200 cc.

Non-Irrititating Iodine Antiseptic

lodine	2 g.
Potassium Iodide	2.4 g.
Alcohol	55 g.
Water	45 g .

Tattoo Mark, Removing

First the skin is vigorously rubbed until the outer epidermis comes off; then a paste of quicklinic, just slacked, to which pulverized phosphorus (two tablespoonfuls to a pint) is added and thoroughly mixed, is applied to the tattooed surface and held by a bandage, which is taken off two days later. The crust is left to dry and then fall off itself; in about fifteen days. A second application should be made; a third is rarely necessary. Thus treated, the tattooing disappears completely without the least scar.

Mechanics Hand Protective Coating U. S. Patent 2.021.131

Water	1600	oz.
Sodium Stearate	288	oz.
Glycerin	1155	oz.
Bodium Silicate	906	OZ.
Lemenone	1	0Z.

Volatile Anæsthetics Formula No. 1

Methyl and Ethyl Chloride equal parts

No. 2	
Ethyl Chloride	60 cc.
Methyl Chloride	35 cc.
Ethyl Bromide	5 cc.
No. 3	

Methyl Chloride Ethyl Chloride equal parts Chloroform

An anæsthetic for external use containing

1/2 fl. dr. Chloroform 21/2 fl. dr. Ether 21/2 fl. dr. Liquid Paraffin

is employed when light anæsthesia is required in painful wound dressings or for short operations.

Anæsthesia Chloroform Preservative Add 1% of absolute alcohol and keep in a cool place away from direct light.

X-Ray Contrast Media

1. Barium diet for stomach and intestinal examinations. Boil together

Corn Starch	15 g.
Sugar	15 g.
Cocoa	20 g.
Barium Sulphate	150 g.
Water	500 cc.

2. Barium diet for diagnosis of stenosis of the small intestine.

Barium Sulphate 200 ec. Thick Gruel

3. Barium suppository for rectum examination.

Corn Starch 750 to 1000 cc. Water Boil together and add a suspension of 200 g. Barium Sulphate Water 500 cc.

Cystographic Medium U. S. Patent 1,935,661

Five to 8 per cent aqueous solutions of sodium (or potassium) bismuth tartrate or citrate (1) serves as cystographic media opaque to X-rays; (1) should contain about 65 (70) per cent of bismuth.

> Hormone Manufacture U. S. Patent 1,978,297

The ground whole testicles are preferably macerated from 12 to 48 hours with

the required amount of the solvent selected, the liquid is filtered off, the residue expressed and re-extracted with preferably the same solvent, this time the glands having been freed from the water therein) using the exact concentration which recovers most of the hormone with the least undesired material, as, for example, 90% acctone, 70% propyl alco-hol or about 75% ethyl or methyl alcohol, etc. Extraction is continued until the residue is fully extracted. The extracts are combined, and the solvent distilled off at a low temperature and under reduced pressure. All traces of the solvent are removed, leaving the lipoid material containing the hormone, together with other substances emulsified in an aqueous solution.

The mixture resulting from the agitation of the emulsified aqueous solution of the lipoid material with one of the solvents named above, when the agitation has ceased, separates into two or three layers, dependent upon the solvent used. When three layers are formed, the upper or solvent layer contains the active lipoid with possible traces of cholesterol and phospholipins, and is free from protein, the middle layer contains most of the phospholipins and cholesterol present in the original extract, together with other organic material, and a portion of the solvent and water. The lower aqueous layer contains blood pigments, salts, etc. The one or two lower layers are preferably drawn off and the agitation with the hormone solvent repeated several times and finally the two or three layers are drawn off separately. In case chloroform is used the lower chloroform layer contains the active hormones.

The combined upper layers may then be washed with a 1 to 10% sodium carbonate solution to remove all traces of the fatty acids and phospholipins, washed with water to remove the sodium carbonate and the solvent distilled off. The residue then contains the testicular hormone in a high state of purity.

For example, in using amyl alcohol at this step of the process, the agitated mixture of the amyl alcohol and the aqueous solution containing the lipoid material separates into three layers, with the upper layer containing the active portion or hormone. The two lower layers are then drawn off, the agitation with amyl alcohol repeated and the upper layers resulting from several repetitions of this step combined, washed with a 1 to 10% sodium carbonate solution and then with water and the solvent distilled off leaving the hormone in a high state of purity.

b.

Analgesic Chaulmoogra Oil Chaulmoogra Oil	for Injection 80 cc.
Olive Oil	20 cc.
Benzyl Ephedrine Base	0.1 g.
Intravenous Colloidal	
British Patent 433	,833
Sodium Sulphide, Pure Water, Distilled and	23.5 g.
Deaerated	50 cc.
	• 0
Dextrin	10 g.
Dissolved in	
Water, Distilled	400 cc.
Dilute to	1 1.
Add sulphur dioxide to a and dilute with distilled wat	n pH of 7.6 er to 10 mg.
of sulphur per cc.	

Hydrogen Peroxide Preservative

The addition of 20 g. phenacetin to 5 kg. hydrogen peroxide acts as a good preservative.

Preservatives for Hydrogen Peroxide

According to French chemists the best preservative for hydrogen peroxide solution is phenetidine lactate in the proportion of 0.5 g. per liter of solution. Less effective are glucose, gelatin (0.2 g. per liter); ethyl alcohol (16 g. per liter); and hippuric acid (0.2%).

Embalming Fluid-For Decolorizing Jaundice Cases

U. S. Patent 1,942,407

Benzoyl Peroxide	15	g.
Ethyl Alcohol (95%)	3	gal.
Formalin (40%)	4	pt.
Water	. 11/2	gal.

Embalming Fluid

Formula No. 1

Formalin (40%)	٠	220	02.
Glycerin		100	υZ.
Borax		90	0 Z .
Sodium Chloride		10	oz.
Sodium Nitrate		10	oz.
Potassium Citrate		50	0 z .
Methanol		40	oz.
Water		75	oz.
Benzaldehyde		6	oz.
Color with Erythrosina			

No. 2

Borax	4	oz.
Phenol	5	0Z.
Salicylic Acid	5	0 z. "
Formalin (40%)	71	0Z.
Glycerin	31	0 Z.
Water sufficient to	make 1	oal.

Corpse Wound Filler	
Yellow Beeswax	5 oz.
Paraffin	5 oz.
White Petrolatum	15 oz.
Sonp Flakes	2 oz.
Water	5 oz.

Finishing Cream	(Corpse)	
Glycol Stearate	12	OZ.
Glyceryl Tristearate	5	oz.
Rose Oil	2	OZ.
Glycerin		OZ.
Water		OZ.
Titanium Dioxide	1	OZ.

Animal Embalming Fluid

Use a water solution of either 5% furfural or 10% formaldehyde.

Air Purifier

Alcohol (95%)	2000	cc.
Formalin (40%)	400	cc.
Pine Needle Oil	190	cc.
Thyme Oil	10	cc.
For use dilute with water	1:50.	

Solid, Volatile Preparations to Perfume and Disinfect the Air

> Formula No. 1 Naphthalene, Pure

Paradichlorobenzol, Pure

No. 3*

Naphthalene, Scales	70 g.
Camphor, Sublimed	10 g.
Paradichlorobenzol	20 g.
No. 4	

No. 4

Naphthal	ene		80	g.
	Acid (Phe	nol)	20	
Heat the	mixtures	gently.	verv	lit

beyond the melting point (color optionally with yellow, red, blue, oil soluble dyestuffs) and pour into molds. Work in well ventilated rooms.

* 0.5% of Citral may be added.

Water Soluble Bactericide U. S. Patent 1,930,474

A 1:1 mixture (200 g.) of chloro-thymol and olive oil is treated with sul-phuric acid (60 g.) at 20° for 2 days, and then washed free from acid with saturated aq. sodium sulphate; the prod-uct is readily dispersed in water.

Protecting Tin Collapsible Tubes Against Corrosion

U. S. Patent 1,968,722

Collapsible tubes containing soap, shaving cream, toothpaste and other alkaline materials are protected against corrosion by addition of 0.1% sodium nitrite.

Pharmaceutical Charcoal Preparations Tablets

Formula No. 1	
Activated Carbon	200 g.
Gum Tragacanth	8 g.
Sugar	195 g.
Water	68 g.
No. 2	-
Activated Carbon	100 g.
Sugar	5 g.
Albumen Solution	5 g.
Gum Arabic	3 g.
Tincture of Benzoin	1 g.

The above are useful in the treatment of dyspepsia.

Removing Creosote from Skin and Clothing

Wash with isopropyl alcohol to remove crossote and prevent further "burning" of skin.

Zinc Ointment

White Beeswax	60 g.
Spermaceti	60 g.
Oil of Sweet Almonds	300 g.
Digest 2 hours on water	bath.
Gum Benzoin, Siam	20 g.
Add while cooling.	_
Zine Oxide	100 g.
Boric Acid	2 g.
Carmine enough	ı to color
Perfume with extract of ro	se leaves.

Hiccough Remedy

Take one teaspoonful of tincture of castoreum and repeat in a half hour if needed.

Fingernail Cleaner

A fingernail stain remover consists of a saturated solution of tartaric acid in water.

EMULSIONS

GASOLINE EMULSIONS

		GASOLINE EM	ULSIONS		
Formula N	o. 1	Oleic Acid	1 cc.	No. 17	7
Tricthanolamine	11/2 cc.	Alcohol	3 cc.	% C. Wa	
Water	1 cc.	Gasoline	45 cc.		
Oleic Acid	11/2 ec.	No. 9		Triethanolamine	175 cc.
Butanol	5 cc.	Triethanolamine	1 cc.	Water	260 cc.
Gasoline	45 cc.	Water	1 cc.	Stearic Acid	1400 ce.
		Steame Acid	5 cc.	Alcohol	1100 ce.
Dissolve trieth		Alcohol	3 cc.	Gasoline	31500 cc.
in water and add ture of other in		Gasoline	45 cc.	No. 18	1
slowly while stirr		No. 10	10 (()	1/2% Wa	ter
ously.*	ing vigor-	Triethanolamme	1 ec.	Triethanolamine	175 cc.
•		Water	1 cc.	Water	175 cc.
No. 2		Stearie Acid	3 ce.	Steame Acid	1400 cc.
Triethanolamine	1 cc.	Alcohol	3 cc.	Alcohol	1400 cc.
Water	1 cc.	Gasoline	45 cc.	Gasoline	31500 cc.
Oleic Acid	1 cc.	No. 11	40 ((,		
Butanol	5 cc.			No. 19	
Gasoline	45 cc.	Triethanolamine	1 cc.	1/1% Wa	ter
No. 3		Water	1 cc.	Triethanolamine	175 cc.
Triethanolamine	1/2 cc.	Stearie Acid Alcohol Gogolina	2 cc.	Water	85 cc.
Water	1 cc.	Alcohol	3 cc.	Steame Acid	1400 cc.
Oleic Acid	¼ ec.	Gasonno	45 cc.	Alcohol	1400 ce.
Butanol	5 cc.	No. 12		Gasoline	31500 сс.
Gasoline	45 cc.	Tricthanolamine	1 cc.		
	10 (1.	Water	1 cc.	No. 20	
No. 4		Stearic Acid	3 cc.	10% Wa	ter
Triethanolamine	1/2 cc.	Alcohol	2 cc.	Trihydroxyethyl-	
Oleic Acid	1 cc.	Gasoline	45 cc.	amme Laurate	3500 cc.
Water	1 cc.	No. 13		Gasoline	31500 ec.
Butanol	5 cc.	Triethanolamine	1/2 cc.	Butanol Water	5600 cc.
Gasoline	45 cc.	Water	1 cc.	Water	3500 cc.
No. 5		Stearic Acid	3 ec.	Triethanolamine	1750 ec.
Triethanolamine	1 cc.	Alcohol	3 cc.	No. 21	
Water	1 cc.	Gasoline	45 cc.		
Oleic Acid	1 cc.	No. 14		5% Wat	er
Butanol	2 cc.	Triethanolamine	1/4 cc.	Trihydroxyethyl	
Alcohol	4 cc.	Water	1/2 cc.	amine Laurate	
No. 6		Steame Acid	3 cc.	Chasoline	31500 cc.
Triethanolamine	1 cc.	Alcohol	3 cc.	Butanol	3500 cc.
Water	1 cc.	Gasoline	45 cc.	Water	1750 ec.
Oleic Acid	1 cc.	No. 15		Triethanolamine	1050 ec.
Butanol	4 cc.	Triethanolamine	1/4 cc.	No. 22	
Alcohol	2 cc.	Water	1/2 cc.	1% Wate	o r
	= 00.	Stearic Acid	2 cc.	,	
No. 7	_	Alcohol	2 cc.	Trihydroxyethyl- amine Linoleat	o 9100 aa
Triethanolamine	1 cc.	Gasoline	45 cc.		31500 cc.
Water	1 cc.	No. 16		Butanol	3500 cc.
Oleic Acid	1 cc.			Water	700 ec.
Butanol	5 cc.	1% Wate		Tricthanolamine	
Kerosene	20 cc. 25 cc.	Triethanolamine	175 cc.	11 10 CHAHOLAMING	2000 004
Gasoline	20 cc.	Water	350 cc.		
No. 8			1400 cc.		
Triethanolamine	1 cc.	****	1400 cc.		
Water	1 cc.		31500 сс.		
* The stability of	the above	emulsions is improve	d considera	bly if they are pass	ed through

^{*}The stability of the above emulsions is improved considerably if they are passed through a colloid mill.

aп

Bright 1	Drying Wax En	nulsion
Paraffin W	ax	15 g.
Oleic Acid		15 g.
Triethanola	mine	20 g.
Borax)	previously	
Water }	dissolved	71/2 g.
electric mixe	er to 90° C. an er. While keep	ing at 90-
	rring vigorousl	
lowing which	must be at 90-	-95° C.

100 foll 100 g. Carnauba Wax Water 1000 cc.

Cool quickly and package.

Paraffin Wax Emulsion Formula No. 1

120 g. Paraffin Wax Stearic Acid 12 g. Melt together and while stirring vigorously add following heated to 55° C. Ammonia (26° Bé.) 6 cc. Water 182 cc. Stir until uniform. No. 2

Glyceryl Monostearate 5 g. Water 150 cc.

Heat and stir vigorously until uniform. Pour into this slowly while stirring strongly: 40 g.

Paraffin Wax (melted)

Paraffin Wax Emulsion (Non-Alkaline)

25 g. Paraffin Wax Glycol Stearate 5 g. Melt together and while stirring vigorously add Water (boiling) 175 cc.

Laundry Calendering Wax Emulsion Mix 33 parts of paraffin wax with 3 arts of oleine, and pour this mixture into a solution of 0.6 part of strong ammonia in 63.4 parts of water, heated to 160° F.

Aqueous Fat-Dissolving Emulsion German Patent 598,216

Prepare: Carragheen Moss Dispersion, wasning gently in water, remove, thicken components in a centrifugal.

Acidify with oxalic acid. Mix thoroughly.

Acidified Carragheen Moss Solution 100 cc. Phosphoric Acid (Free from 10 cc. Arsenic) (67%)

200 cc. Fat Solvent e.g. Trichloroethylene 100 cc. Naphtha 200 сс.

Chlorinated Naphthalene Emulsion British Patent 413,756

Eighty g. of wax-like chlorinated naphthalene, of setting point 93° C. is dissolved in 100 g. trichloroethylene, and is added with stirring to a warm mixture of water 60 g., Turkey-red oil 10 g., casein 3 g. and strong ammonium hydroxide 1 g.

Emulsions of Oils, Fats, Waxes and Resins

British Patent 431,642

Water is dispersed in oils, fats, waxes, resins, artificial resins, pitches, asphalts or the like by adding to the water, prior to or during the mixing, about 0.01% of the principal substance of aqueous alkali, such as caustic soda or potash or ammonia, having dissolved therein aromatic hydrocarbon derivatives or their salts soluble in alkali such as benzoic acid, sodium salicylate, o, m or p-cresol, xylenol, guaicol, or cresol, or mixtures thereof. The products may have pig-ments or solid substances incorporated therewith for use as paints, color varnishes, printing inks, or lubricants.

Formula No. 1

600 g. of water, containing 0.012 g. of a solution of 15% caustic soda and 1.5% of the above specified substances, are stirred at 25-30° C. into 1000 g. of linseed-oil varnish; 280 g. of the resulting water-in-oil emulsion are stirred with 530 g. of red lead and 175 g. of calcite.

250 g. of water containing 0.02 g. of emulsifier as in (1) are stirred into a mixture of 100 g. of asphalt and 900 g. of printers' linseed-oil varnish; 9 g. of nigrosin are stirred into the product to produce a printers' ink.

No. 3

350 g. of water containing 0.015 g. of caustic soda and 0.0015 g. of sodium benzoate are stirred at 30° C. into 1000 g. of olive oil; the product may be used as salad oil.

No. 4

800 g. of water containing 0.02 g. of caustic potash and 0.02 g. of cresol are stirred into 1000 g. of viscous mineral lubricating oil and 100-200 g. of

graphite add	ed with	stirring	to	produce	a
lubricant.					

Emulsions of Oils, Fats, or Waxes German Patent 575,922

Formula No. 1		
Cod Liver Oil	80	g
Pectin	0.5	
Milk Sugar	20	g
Water	20	g
No. 2		_
Paraffin Oil	80	g
Pectin	0,5	g
Milk Sugar	20	g.
Water	20	g.
German Patent 585,586, Ad the Above (575,922		to

For stable emulsions containing up to 80% of oils use instead of Milk-Sugar:

Fruit Sugar (Fructose)
Invert Sugar (Invertose)
Grape Sugar (Glucose)
Manna Sugar (Mannose)
Never use Cane Sugar!

Pine Oil Emulsion

Pine Oil	9	g.
Diglycol Laurate	4	g.
Mineral Oil	1	g.
Water	100	g.
Mir fret three meterials	and than	9.

Mix first three materials, and then add water slowly while stirring vigorously.

Cottonseed Oil Emulsion

Diglycol Laurate	18 cc.
Cottonseed Oil	40 cc.
Water	50 cc.

China Wood Oil Emulsion

Omme 1100a	0.12 22.00	
Diglycol Laurate	18	ec.
China Wood Oil	40	ee.
Water	55	rc.

Mineral Oil Emulsion Cream

Glyceryl Monostearate Water	$\frac{5}{125}$	g. cc.
--------------------------------	-----------------	-----------

Heat together and stir until uniform then add slowly while stirring vigorously Mineral Oil 43 cc.

Soluble Oil

U. S. Patent 1,965,935

A soluble oil composed of the following ingredients has unique emulsification and stability properties:

	4 g. 6 g.

Mineral Oil	64	g.
Water White Rosin	10	g.
"Carbitol" (Monoethyl		
Ether of Diethylene Glycol)	2	g.
Diethylene Glycol	4	g.

Diethylene Glycol 4 g.

This oil is clear and will not become cloudy when cooled to a temperature of 60° F, and will not become covered with a film after standing exposed to the air at a temperature of 80° F, over a long period of time or at a temperature of 200° F, for one day. This oil will readily emulsify with water after standing exposed to the air at 200° F, for two days. Aqueous emulsions containing this oil are very stable even at a temperature of 200° F. In general, stable aqueous emulsions are prepared by using 1% to 35% of this oil, although stable aqueous emulsions can be prepared by using proportions of the oil outside these limits.

Carbon Tetrachloride and Tetrachloroethylene Emulsions

The following formula may be used for a 50% preparation: 20 g, of connectial soft soap, 6 cc, of cresol, 50 cc, of carbon tetrachloride or tetrachloroethylene and 100 cc, of liquid parafilm.

Phenol-Formaldehyde Resin Emulsion Australian Patent 17,583

Phenol Formaldehyde Resin	45	g.
Paraffin Oil	5	g.
(Heat together)		
Sulphonated Sperm Oil	5	g.
Olein	5	g.
Cyclohexanol	1	g.
Partially saponify above wi	th R	meons

Partially saponify above with aqueous caustic soda then add

Glue 2½ g. Water 45 g. Mix in homogenizer or colloid mill.

Synthetic Resin Emulsion U. S. Patent 1,999,715

One hundred parts, by weight, of ground resoremol are placed in a metal container or kettle which is jacketed with an outside containes in turn equipped for steam heating and water cooling. This permits of temperature regulation during the chemical reaction as it is very essential to carefully control the temperature of the reacting substances in order to prevent their conversion to the insoluble

stage. The kettle may be equipped with suitable agitators to permit the rapid churning of the kettle contents. To the resoremed, 112 parts, by weight, of 37.5% formaldehyde solution (formalin) are added, and the temperature is increased to a maximum of 60° C., so that the resoreinol will dissolve.

In a separate container, 8 parts, by weight, of para-nitraniline are dissolved in 20 parts, by weight, of cresol having a boiling point range of from about 215° C. to 230° C. The melted paranitraniline is added to the formaldehyde solution, and the mass is thoroughly agitated, the reaction temperature being raised to from 70° C. to about 75° C. A plasticizer of vegetable or animal oils and wax with a filler and suitable coloring material may be added as a paste to the mixture in the kettle. As an example, employ 8 parts, by weight, of chay, 0.8 part, by weight, of beeswax, and 1 part, by weight, of iron oxide, all best ground in a paint mill or ball mill to obtain thorough dispersion. The paste has been found to mix readily with the thickening liquid in the kettle.

For best results, the temperature should be maintained at no higher than 80° C. When the mass in the kettle becomes stringy, and before gelatinization can take place, the emulsification cahancing agent is added. A water solution (65 parts, by weight) containing 0.1% of borax is added, first slowly, and then rapidly to the agitated resin. The borax, or any other suitable alkaline salt, serves to so enhance the emulsifying action of the wax and/or the oils (and gnms, if any are used) as to enable them to properly maintain the resin in suspension in the water. This results from the fact that the borax or other alkaline salt used reduces the surface tension of the water from its normal value at the operating temperatures, thereby more readily effecting a wetting of the resin particles and embling the wax or other fatty material used to more easily retain the resin particles in suspension. The temperature is dropped to about 20° C. and thickening of the resin takes place. A solution of alcohol (about 65 parts, by weight) or an equal quantity of a 20% bengol or toluene solution in alcohol may be added to the emulsifier to obtain proper consistency. The varnish is immediately strained to remove any foreign particles. The mass is then cooled, with accompanying increase in viscosity of the varnish, and additional alcohol or solvent mixture may be added to obtain the desired viscosity.

Chlorinated Rubber (Tornesit) Emulsions

U. S. Patent 2,008,558

Formula No. 1

50 parts, by weight, of toluene, 50 parts of water and 20 parts of pulverulent chlorinated rubber are introduced jointly into a stirring apparatus and stirred. A uniform and stable dispersion is formed in a few minutes.

No. 2

50 parts, by weight, of toluene and 50 of water are brought together and intimately stirred, 20 parts of finely divided solid chlormated rubber being added during the stirring operation. A uniform and stable dispersion is formed immediately.

Chlorinated Rubber Emulsion British Patent 414,072

20 parts, by weight, of oleic acid, saponified with 20 parts of sodium silicate in 200 parts of water, is stirred at 100° C. into 5 parts of chlornated rubber dissolved in 25 parts resin oil; 125 parts of water containing casein 8 and ammonia 0.5 parts is then added.

Aqueous Dispersions of Bitumen German Patent 557,228

a. Soya Bean Meal Water	1 g.
". Water	49 cc.
b. Caustic Soda	0.2 cc.
o. Bitumen Mixture,	
Liquefied	50 g.

Boil a after sonking, then saponify with b and emulsify c, stirring vigorously.

Tar Emulsion

Austrian Patent 137,89	1	
Crude Montan Wax Crude Wool Fat Tar	2	lb. lb. lb.

Heat to 80-90° C. and while mixing vigorously run into a 1% caustic soda solution heated to 60° C.

Asphalt Emulsion U. S. Patent 1,931,072

An aqueous solution of soap (9 parts) by weight, is dissolved in warm water (78 parts), and a low grade fuel oil or crude asphaltic-base oil (20 parts) is dispersed therein. A relatively small quantity (1-2 parts) of a metallic salt

100 g.

of a fatty acid, e.g., aluminum oleate is mixed therewith, the emulsion is warmed, and asphalt (296 parts) is added slowly and with agitation, and distributed uniformly throughout the mixture.

Non-Rusting Alkaline Emulsions Latherless Shaving Cream U. S. Patent 1,968,722

Stearic Acid	22	lb.
Glycerin	10	lb.
Ammonia (28%)	1	lb.
Sodium Nitrite	0.1	lb.
Water	67	lb.
The stearic acid is first hea	ted to	about
5° C. The glycerin and	wate	r are
in 1 Amethor appri	from	the

re then mixed together apart from stearic acid and also heated to about 85° C. To the glycerin and water add the ammonia. This solution is then poured into the stearic acid and thoroughly stirred. When the whole mix is cooled add sodium nitrite.

Polish

Carnauba Wax	12 g.
Rosin	0.5 g.
Triethanolamine Oleate	3.5 g.
Sodium Nitrite	0.1 g.
Water sufficient to make	e 100 g.

Library Pasto

Starch	24 g.
Gum Acacia	3 g.
Glycerin	6 g.
Borax	0.5 g.
Sodium Nitrite	0.1 g.
Oil of Cloves	0.1 g.
Water	72 g.
Soap Base Lubricating	Emulsion

3 kg. Cottonseed Oil 1-2 kg. Mineral Oil 132 g.

Caustic Soda

Heat to 180° C. until foaming stops. Add 13 kg. mineral oil in successive portions at intervals with stirring bringing up to 190-210° C. Pour into wooden tubs and cool to 70° C. Add 9 kg. water with stirring.

High-Molecular Organic Sulphonic Acid Emulsifier

German Patent 616,321

Formula No. 1

Yellow Oil from Brown	
Coal Tar	100 g.
Paraldehyde	10 g.
Chlorosulphonic Acid	125 g.
Add the acid at 30-35° C.	, cool, st

ir

sulphonic acid from unchanged oil. Pour into 3 parts of ice-water neutralized with concentrate caustic soda. Let stand, separate from impurities, dry in vacuum.

No. 2

Same, but substitute Paraldehyde by Acetalde- hyde (50%)	20 g.
No. 3	
As No. 1, but use	

Paraffin Oil from Brown Coal Tar

fact as in No. 1.

No. 4

Solar Oil from Brown Coal Tar 20 g. Heptaldchyde (Oenanthol) Chlorosulphomic Acid 150 g. At 35° C. has to stand 1 day, other

No. 5

Paraffin Oil (7.5° E at	
20)	100 g.
Benzaldehyde	15 g.
Chlorosulphonic Acid	130 g.
Mathod on No. 1.	

Sulphonation of Cetyl Alcohol

Melt the Cetyl Alcohol	40	g.
Dissolve in Acetic Anhydride	20	g٠
Treat with Sulphuric Acid (Concentrated or Funning) The reaction is run below 10°	40 C.	g.

Sulphonating Napthenic Alcohols U. S. Patent 2,000,994

One part by weight of a raw commerone part by weight of a raw commer-cial naphthenic acid (boiling point 90-230° C, at 13 mm. pressure) is dissolved in 2 parts by weight of 3% butyl alco-holic hydrochloric neid and heated to boiling for four hours. The butanol and hydrochloric acid are then distilled off and 200 kg. of the naphthenic acid so treated are reduced in an autoclave with 90 kg. of sodium and 1,000 kg. of butyl alcohol. The whole is then heated under constant agitation to 140° C. for 11/2 hours. After cooling to 90° C. the reaction mass is poured into water, the underlying liquor is drawn off and the remainder is neutralized and washed several times. It is then dried over lime and the excess butyl alcohol is removed by distillation. The product so obtained boils between 70 and 230° C. at 10 mm. thoroughly. After 18 hours separate the pressure and posseses an acetyl saponification number 175 and an iodine number 22. It is free from saponifiable components and dissolves to give a clear solution in concentrated sulphuric acid. Dilution with water produces no turbidity. The conversion of the product into the sulphuric acid derivative can be carried out in the following manner:

20 parts by weight of chlorosulphonic acid are gradually added to 50 parts of the above mentioned product and to this are subsequently added 5 parts of sulphuric acid whereupon the temperature rises to 40° C. The reaction mass is then washed with salt solution and neutralized. Upon evaporation in vacuo the sulphonate and/or sulphate is obtained in a solid grindable form.

Sulphonating Oils

A. Cod, Sperm, Cottonseed and Castor Oils

1. High Sulphonation Product

Any of the Above Oils 735 lb. Sulphuric Acid 275 lb.

Run in the acid in a thin jet as quickly as possible with good mixing but do not allow temperature to rise above 95° F. Agitate for 5 or 6 hours until a sample in the case of cod oil is soluble in distilled water without opalescence. With cottonseed oil the solution will be slightly translucent. The oil is now dropped into the mixing tank, containing two and onehalf times the volume of oil of Glauber's salt solution, 10° B6. Agitate smoothly for five to ten minutes and warm to 104° F. Allow to separate. Draw off the water and make the oil nearly neutral to methyl orange with caustic soda. Allow to stand over night. It is to be noted, that according to the acidity of the oil at this stage, when allowed to stand, the amount of free fatty acid in the finished oil can be regulated. Next morning, draw off the water again, and clear with caustic sods.

B. Red, Cod, Castor, Neatsfoot or Refined Corn Oils

2. Quick Sulphonation Method

Any of the Above Oils 775 lb. Sulphuric Acid 225 lb.

Usually used for oleic acid, cod oil, castor oil, Neatsfoot oil, refined corn oil, and mixed oils. Sulphuric acid = 22½% on the weight of the oil.

on the weight of the oil.

The acid is run into the oil quickly while the oil is violently agitated. With a ten-barrel batch of oil, the acid takes about thirty minutes to run in. The temperature rises quickly and as soon as it

reaches 130-135° F., the oil mixture is dumped quickly into a mixing tank situated underneath the sulphonating tank. The mixing tank contains Glaub er's salt solution 10° Bé, equal to double the volume of the oil. The Glauber's salt solution is at room temperature. The oil and Glauber's salt solution is agitated smoothly for five to ten minutes and the oil allowed to separate. Separation is nearly complete in half to one hour. The clear water is drawn off to a storage tank, and after neutralizing with caustic soda, is used over again for the next batch. The oil is neutralized with caustic soda until it is nearly neutral to methyl orange, that is, slightly on the acid side. Allow the oil to stand until morning and a further separation will take place. When the oil is completely separated, and the water drawn off, the oil should test 20% water. It is now cleared by the addition of further caustic soda. In winter time, it is better to use caustic potash for the final finishing, as it gives a more liquid oil. In testing the acidity of the oil, after the first separation, it is recommended to use an ether and salt solution for the titration with methyl orange.

C. Castor Oil, Concentrated

Castor Oil 1000 lb. Sulphuric Acid (100%) 1000 lb.

Dilute the castor oil with ethylene dichloride. Run the acid in slowly to the previously cooled oil and solvent mixture. Do not allow the temperature to rise above 60° F. After the acid is all in, continue stirring until a few drops dissolve perfectly clear in distilled water. and also dissolve perfectly clear in a saturated solution of calcium sulphate. Do not continue stirring after this point, but then add to it a 5% solution of Glauber's salt solution, equal in volume to three times that of the sulphonated mixture. The solution of Glauber's salt is kept cool by means of ice. The temperature not being allowed to rise above 60° F. Allow to separate, and wash twice with 25% Glauber's salt solution. Separate, and add caustic soda until neutral, and then distil off the solvent.

D. Oleic Acid and Ricinoleic Acids, Concentrated

Above Fatty Acids 100 lb. Sulphuric Acid (100%) 100 lb.

On a large scale some manufacturers use a dough type mixer, brine cooled, while others use a system, wherein the sulphuric acid and oil are sprayed simultaneously by a whirl disc system into a large reaction vessel, being sufficiently cooled previously so that the heat of reaction does not cause the product formed to become unduly heated before running out of reaction vessel. Sulphonation uses 100 lb. fatty acids and 100 lb. sulphuric acid 100 per cent strength.

Keep temperature below 50° F. while adding the sulphuric acid. Sulphonation time is 50-60 minutes. Wash with Glauber's salt solution 12-15° Bê. twice, keeping temperature below 70° F. Let stand over night to separate and neutralize with caustic soda. The product is allowed to stand 3-5 days at 15-20° C. to allow the Glauber's salt to crystallize out. This crystallization can be improved by the addition to the oil of a small quantity of a volatile solvent such as xylene, trichloroethylene, carbon tetrachloride, etc.

Sulphonation of Castor Oil French Patent 745,787

a. Castor Oil 100 kg. b. Sulphuric Acid (66° Bé.) 100 kg.

Add b to a in very thin jet (2 hours) and with continuous stirring, keeping the temperature at $10-13^\circ$ C. Wash product once or twice with salt-solution, keeping the temperature below 15° C. Separate oil from the aqueous layer in the usual way, neutralize.

Sulphonating Oils U. S. Patent 1,967,655

Formula No. 1

100 kg. of ricinoleic acid are sulphonated at temperatures below 5° C. with 90 kg. of 30% oleum. 30 kg. of glycol mono-methyl ether are added to the crude sulphonation product. After completion of the reaction, ice is added and the product washed with Glauber's sult solution.

No. 2

100 kg. of 12-hydroxy stearic acid are mixed with 65 kg. of glycol mono-ethyl ether and sulphonation effected at temperatures below 0° C. with 36 kg. of chlorosulphonic acid. The product is worked up as in No. 1.

No. 3

100 kg. of naphthoic acid are sulphonated with 70 kg. of chlorosulphonic acid, 55 kg. of glycol mono-methyl ether are added to the crude sulphonation product. In place of sulphuric acid and the like sulphonating agents, alkyl sulphuric acids

or alkyl chlorosulphonic acids may be employed.

Sulphonation of Fatty Oils, Fats, Waxes
Austrian Patent 134,993

Whale Oil (Sperm Oil) 1 lb.
Spindle Oil 3-4 lb.
Funning Sulphuric Acid

(30% SO₃)

Run reaction at 40-45° C., adding sulphuric neid in a jet. Stir, then let stund 12 to 24 hours; wash with sodium chloride or sodium sulphate-solution, separate from acid wash water. Neutralize, if necessary, with organic bases, until a drop of oil, when diluted with water,

Cellulose Ester Emulsions U. S. Patent 1,970,572

shows nearly no turbidity.

A pyroxylin base is prepared by colloiding 12.5 parts by weight of alcohol-wetted pyroxylin (10 parts of dry \(\frac{\psi}{n}\) pyroxylin) with 20 parts by weight of blown lasced oil in a suitable mixer, such as the Werner and Pfleiderer mixer. 25 parts by weight of a solvent mixture are then added to the colloided mass in portions equalling 5 parts by weight to form a homogeneous base having the following composition:

Pyroxylin (1/2 sec.)	10 g.
Alcohol (Denatured)	2.5 g.
Blown Linseed Oil	20 g.
Butyl Acctate	20 g.
Butyl Lactate	5 g.

An emulsion is prepared by heating 0.5 part by weight of sodium olente with 15 parts by weight of gasoline to a clear gel; after which 2 parts by weight of water are added to the hot gel with vigorous stirring, thus forming a concentrated emulsion of gasoline in water that is stabilized by sodium oleate. For convenience this will hereafter be called the agent emulsion.

The presolution or solvating of the sodium oleate in gasoline or some similar liquid is desirable to assure uniform distribution.

17.5 parts by weight of the agent emilsion are then stirred vigorously into 57.5 parts by weight of the pyroxylin base with a high speed stirrer of the propeller blade type.

propeller blade type. Inversion of the emulsion from the water-in-oil type to the oil-in-water type may be effected in various ways, as explained below, but in this example it is effected by the sudden addition of water in relatively large quantities, the time of

addition being the controlling factor in particle size, as indicated by systems a.

b, and c.

System (a): 20 parts by weight of water are added in small portions with vigorous stirring, thus yielding a viscous water-in-oil type dispersion. 10 parts by weight of water are then added with vigorous stirring to invert the system to the oil-in-water type. 68 parts by weight of water are added next, either slowly or rapidly, with more moderate stirring. Microscopic measurements of particle size average 1.19 microus, and the dispersion spontaneously wets an absorbent type of paper.

System (b): 35 parts by weight of water are added in small portions with vigorous stirring, yielding a viscous water-in-oil type dispersion. 10 parts by weight of water are then added with vigorous stirring to invert the system to the oil-in-water type. 53 parts by weight of water are next added, either slowly or rapidly, with more moderate stirring. The average particle size is 1.92 microns, and the dispersion does not wet paper spontaneously. Vacuum filtration is re-quired in order to effect paper penetration, and some separation of disperse

phase occurs on the surface of the paper.

System (c): 90 parts by weight of
water are added in small portions with vigorous stirring, yielding a viscous water-in-oil type dispersion. 8 parts by weight of water are then added with vigorous stirring to invert the system to the oil-in-water type. The average particle size is 2.23 microns. Severe separation of the disperse phase occurs on the surface of the paper during vacuum filtration, and the dispersion is not adapted to paper impregnation.

Petroleum Demulsifier

Diglycol Laurate	83 g	۲.
Sodium Silicate	5 g	ġ.
Rosin Soap	5 g	
Phenol	4 2	•
Water	11/2 g	
Paraffin	1½ g	ζ.
	0	-

Margarine Emulsifler

Refined and deodorized sunflower oil oxidized with a current of dry air at 250° C. for 10 hours shows better emulsifying properties than Paalsgaard oil. When 0.4%, of this oil is added to the emulsified mixture in the manufacture of margarine, the product after standing 54 hours retains the good taste and odor and high moisture content (14.6%).

Breaking Petroleum Emulsions U. S. Patent 1.976.602

React 250 lb. of phthalic anhydride with 500 lb. of castor oil at a temperature of approximately 120 to 145° C. for approximately 6 to 12 hours. The reaction can be followed roughly by withdrawing a small sample of the partially reacted mass and permitting it to cool on a watch glass. When the reaction is completed, crystals of phthalic anhydride no longer appear. When the sample no longer shows the presence of such crystals on cooling, it can be titrated with a standard volumetric alkaline solution, so as to indicate that the acid which remains is due entirely to the carboxylic hydrogen and not due to any unreacted phthalic anhydride. One must guard against a rise in temperature.

The product of reaction represents a viscous yellow oil not unlike blown castor oil in consistency. It is neutralized with sufficient ammonium hydroxide to completely convert all acidic material into the ammonium salt. The product thus obtained is substantially water-soluble

and is suitable for use.

A treating agent or demulsifying agent of the kind described may be brought in contact with the emulsion to be treated in any of the numerous ways now employed in the treatment of petroleum emulsions of the water-in-oil type with chemical demulsifying agents, such for example, as by introducing the treating agent into the well in which the emulsion is produced, introducing the treating agent into a conduit through which the emulsion is stored, or introducing the treating agent into a container that holds a sludge obtained from the bottom of an oil storage tank. In some instances, it may be advisable to introduce the treating agent into a producing well in such a way that it will become mixed with water and oil that are emerging from the surrounding strata, before said water and oil enter the barrel of the well pump or the tubing up through which said water and oil flow to the surface of the ground. After treatment the emulsion is allowed to stand in a quiescent state, usually in a settling tank, at a temperature varying from atmospheric temperature to about 200° F., so as to permit the water or brine to separate from the oil, it being preferable to keep the temperature low enough so as to prevent the valuable constituents of the oil from volatilizing. If desired, the treated emulsion may be acted upon by one or the other of various kinds of apparatus

now used in the operation of breaking petroleum emulsions, such as homogenizers, hay tanks, gun barrels, filters, centrifuges or electrical dehydrators.

The amount of treating agent on the anhydrous basis that is required to break the emulsion may vary from approximately 1 part of treating agent to 500 parts of emulsion, up to a ratio of 1 part of treating agent to 20,000 parts of emulsion, depending upon the type or kind of emulsion being treated. In treating exceptionally refractory emulsions of the kind commonly referred to as "tank

bottoms" or "residual pit oils," the minimum ratio above referred to is often necessary, but in treating fresh emal stons, i.e., emulsions that will yield readily to the action of chemical demulsifying agents, the maximum ratio above mentioned will frequently produce highly satisfactory results. For the average petroleum emulsion of the water-in-oil type a ratio of 1 part of treating agent to 10,000 parts of emulsion will usually be found to produce commercially satisfactory results.

FARM AND GARDEN SPECIALTIES

FARM AND GARDEN SPECIALTIES			
Tree Bands for Caterpillar and Flies Grafting Wax			
Formula No. 1	Formula No. 1		
Rosin Oil 9 g.	(Colophony 350 g.		
". Spindle Oil 20 g.	Beeswax 10 g.		
$b. \begin{cases} \text{Slaked Lime} & 6-9 \text{ g.} \\ \text{Spindle Oil} & 65-62 \text{ g.} \end{cases}$	a. {Pitch 60 g.		
Spindle Oil 65-62 g.	Linseed Oil 25 g. Turpentine, Venice 15 g.		
Add a to b, stir violently to homoge-	b. Methanol 85 g.		
nize. Stir until congealing starts. Allow to set for 24 hours.	Melt up a, then stir until cool, add b.		
	No. 2		
No. 2	Linseed Oil 1 lb.		
Rosin 30 g. Linseed Oil* Varnish 20 g.	Turpentine Oil 4 lb.		
Beeswax, Yellow 2 g.	a. Beeswax 3 lb.		
* Or Rape Seed Oil, or Wool Fat, when	Colophony 9 lb. b. Methanol 1 lb.		
a longer catching period is desired.	Dissolve a cautiously, thin with b.		
The melted and well mixed glue is put	No. 3		
on the bark of the tree; over it put a	1		
ring of cloth, fastened with wire, then put over that again a layer of glue, all	Castor Oil 4 lb. Rosin 5 lb.		
around the stock.	Beeswax 1 lb.		
No. 3	Charcoal % lb.		
Colophony 300 g.	Glyceryl Monostearate 1 lb.		
Linseed Oil Varnish 200 g.	Melt and apply with brush. This ex- cludes air and fung; prevents drying		
Yellow Wax 20 g.	out and doesn't injure live tissues.		
D. A. C. W			
Protecting Mixture for Young Trees Against Game	Bleaching Citrus Fruit Blemish		
Ceresin (58–60° C.) 20 oz.	Navel oranges, badly blemished with		
Spindle Oil, Distilled 60 oz.	souty blotch, are thoroughly cleaned by dipping for ½-1 min. in a solution con-		
Dippel's Animal Oil or	taining 0.25 lb. each of boric acid and		
Carbolineum 20 oz.	chloride of lune.		
Melt up and stir until cold.	-		
Codling Moth Tree Bands	Removing Arsenic Residues from Fruits		
Formula No. 1	Wash with a 1% solution of ammonia or caustic soda.		
Cloth is impregnated with	or caustic soua.		
Beta Naphthol Crude 1 lb.	Preserving Color of Leaves		
Red Engine Oil 1.5 pt.	Immerse leaves in		
Apply at 130-132° F.	Glycerin 5 g.		
No. 2	Copper Sulphate 2 g.		
Beta Naphthol Crude 1 lb.	Water 93 cc.		
Mineral Oil (200-300 sec.) 11/2 pt.	Non-Poisonous Fly-Papers		
Gasoline 1 pt. No. 3	Quassia 16 oz.		
Water 2 gal.	Colocynth 2 oz.		
Ammonia (28%) 2.4 fl. oz.	Long Pepper 4 oz.		
Casein 4 oz.	water to make I gai.		
Mineral Oil, Refined (65-75 sec.) 8 gal	Boil until the decoction is reduced to		
,	l 4 pints; strain; dissolve in the clear		
	110		

liquid 4 oz. of sugar. Dip the absorbent paper in this solution.

Cobalt Fly-Papers

Dissolve cobalt chloride, 1 oz., and Tartar Emetic, 1 d., in 1 gal. of the Quassia decoction (formula above), and dip the paper in the resulting solution.

Fly Catcher	
Colophony (Rosin) G	49 g.
Mineral Oil (Viscosity	
31/2-4° E at 50° C.)	36 g.
Lanolin, Anhydrous	4 g.
Beeswax, Pure	1 g.
Castor Oil	2 g.

Moth Powder

Camphor		4	oz.
Benzoin		1	07
Black Pepper		2	04.
Cedar Sawdust		5	0Z.
Mix after reducing	the	solids	to
oarse powder.			

Roach Eradicating Powder

Sodium Fluoride	60 oz.
Wheat Flour	20 oz.
Corn Starch	12 oz.
Cocoa	8 oz.

The sodium fluoride should be in a finely powdered form and thoroughly mixed and then sifted to make certain of a homogeneous product. This may be made into a paste with a minimum of water and placed in new or used crown caps, allowed to dry and laid in roach infested places. It may also be dusted as a powder. The filled caps, however, can be used over again and are cleaned up more readily than the powder.

Mosquito Spray for Outdoor Gatherings Kerosene Containing Py-

rethrum Extract Equivalent to 1 lb. of Flowers (Analyzing 0.9% Pyrethrins) per Gallon 100 gal. Water 50 gal.

Sodium Laurel Sulphate (Emulsifier)

The emulsifier is first mixed with the water and transferred to the tank. The oil is then run in gradually into the tank with agitators and pump working at full speed. After all the oil has been added the pumping is continued until the entire mixture has passed through the hose and

back into the tank two or three times or until the mixture is thick and homogeneous, showing no free oil on the surface. The finished product is then pumped into drums for storing. This constitutes the stock emulsion. Excessive foaming may be chiminated by dissolving about two or three pounds of wool grease (degras) in the kerosene before emulsifying. Any other suitable apparatus for emulsification can be used.

The cost of preparing the concentrated emulsion is about 23 cents per gallon, based on the present price of pyrethrium, which makes out slightly over 2 cents per spray gallon. When purchased, the stock emulsion costs from 30 to 50 cents per gallon, depending on the quantity ordered.

Directions for Spraying

About half an hour before the gathering takes place the area is completely sprayed with the larvacide diluted 1:10 or 1:12, that is 1 part of larvacide is mixed with 10 or 12 parts of water. The spraying is done with a power sprayer capable of developing a pressure of 100 pounds or more per square inch and equipped with a spray gun. Before mixing with water the concentrated stock larvacide should be well shaken. Also the diluted spray should be frequently stirred or agitated in order to secure uniform distribution throughout the spraying operation. The spray is applied in the form of a fine fog directly to the giass, grounds, tents, trees, shrubs, etc. Then the stream is directed upward so as to saturate the atmosphere with the fog. At no time should a course spray be applied, since it is unnecessary and may injure vegetation. The grounds for about 20 feet outside the area should also be thoroughly fogged, especially when tall grass, shrubs, woodland and other vegetation are present offering a hiding place from which adult female mos quitoes may issue suddenly at dusk in large numbers. If the area has been thoroughly fogged one treatment may suffice for two hours or even the rest of the evening. If mosquitoes become bothersome later in the evening, the area on the outside of the "gathering" grounds should again be fogged, directing the stream primarily upward and towards the ground to be protected. This outside fogging may be repeated again if necessary. On small areas, such as back-yards, private lawns, etc., a knapsack sprayer or bucket pump capable of producing a fog spray, of 10 to 15 feet high, can be used.

Weed Killers Formula No. 1

Poisonous:

Arsenite of Soda (Concen-

trated Solution) 1 gal. 20 gal.

Mix through and sprinkle on vegetation to be exterminated, making application on a bright clear day.

No. 2

Non-Poisonous:

Chlorate of Soda 1 lb. Water 1 gal.

Dissolve chlorate of soda in the water and use this solution without further dilution by sprinkling on vegetation wished exterminated.

Weed Killer British Patent 418,061

Formula No. 1

Ammonium Chloride	83 g.
Copper Sulphate	5 g.
Calcium Carbonate	12 g.
No. 2	
Ammonium Chloride	25 g.
Sodium Nitrate	25 g.
Ferrous Sulphate	50 g.

Ragwort Weed Killer

Use ammonium sulphocyanide 10% solution), 200 gal. per acre. Dry weather is best time.

Killing Weeds on Lawns

To kill weeds on lawns, golf courses, etc., treatment with a solution of ammonium sulphate and soft soap has been found effective. A mixture adopted for this purpose in England contains 1 lb. of ammonium sulphate, 1/2 lb. of soft soap, and I gal. of water, to be used for every 8 square yards.

Hydrogen Sulphide as an Insecticide and Fungicide

Extensive trials carried out by the Azov-Black Sea branch of the All-Union Institute of Plant Protection have proved that hydrogen sulphide may be successfully used for rodents, insects, and fungi control. Laboratory experiments have shown that a 0.02 to 0.03% concentration of hydrogen sulphide in air is sufficient to kill the earless marmot. Field experiments proved that 4 to 6 g. of hydrogen sulphide per burrow are quite sufficient, while in better conditions (i.e., when the

soil is warm and dry) this rate may be reduced to 3 g. per burrow. The same results are obtained when applying sulphuric slags, which emit hydrogen sulphide because of the action of moisture absorbed from the air. In this case some 8 to 9 g. of slag per burrow are sufficient, the mortality of earless marmot reaching 92 to 98%.

Hydrogen sulphide proved especially efficient as a means of destroying barn mites, being more penetrable in grain than chloropicrin and carbon disulphide.

Experiments made in the huge Millerovo elevator have proved the practicability of this method. Exposure for 40 hours at a rate of 400 g. of hydrogen sulphide per ton of grain proved efficient. Hydrogen sulphide does not reduce the germination rate of seeds and only a few strams decrease their germination with 4 to 8%, while the majority of strains even increase their germination rate with 15 to 30 per cent. Experiments on feeding the treated grain to cocks and rabbits

have shown that no injury has resulted. Fair results have been obtained, too, when applying hydrogen sulphide as a fungicide. Laboratory experiments have proved that the spores of main fungous diseases of seeds perish when seeds are exposed to hydrogen sulphide for 1 to 4 days at a rate of 200 to 400 g, of gas per cu. m. (smut and bacteriosis of cereals, gommosis of cotton, bacterial rot of vegetables).

Red Squill Extract

Extract 15 g. red squill by repeated extractions with 100 ec. boiling methanol in an enclosed system percolutor.

Insect Spray Formula No. 1

Petroleum Spirits	1000	cc.
Pyrethrum Extract		g.
Sassafras Oil	5	cc.
Methyl Salicylate	20	cc.
No. 2		
Petroleum Spirits	550	cc.
Vaschne Oil	450	ce.
Methyl Salicylate	20	cc.
Sassafras Oil	10	cc.
Pyrethrum Extract	10	g.

Insecticide Spray Spreader

Water	5 cc.
Caustic Potash	7.4 g.
Pine Tar Oil	44.3 cc.
"Cellosolve"	10 cc.
Oleic Acid	33.3 сс.
Mix in the order given.	

Light Stable Insecticid	e Spre	ıy
U. S. Patent 2,011,	428	
Gum Ghatti	2.4	lb.
Cresylic Acid	0.18	lb.
Water	35	lb.
White Oil (80 sec. Saybolt		
at 100° F.)	62.4	lb.
1,4 Toluido Anthraquinone	0.02	lb.
The concentrated emulsion pared by intimately mixin gredients in a colloid mill or the mixture through a centrior in any other suitable man a concentrated emulsifiable which may readily be diluted emulsion suitable for spraying	g the by pr fugal ner to rompo to yie	e in- assing pump give sition dd an

Codling Moth Control by Nonarsenical Sprays

Sprays containing nicotine sulphate (1:640) and white oil (1:80) gives much better control of the codling moth than lead arsenate sprays.

Non-Poisoning Fruit Spray Formula No. 1 Diglycol Laurate Pyrethrum Extract (20 Fold)

Water	100 gai
No. 2	
Derris Extract (5%)	1 qt.
Skim Milk Powder	1 lb.
Water	100 gal.
No. 3	
Derris Root, Ground	10 lb.

Orange Worm Spray Formula No. 1

90 gal.

Potassium Aluminum Fluoride 50 lb. 45 lb. Fiber Tale Mineral Oil, Refined 5 lb. (70 Viscosity)

Use 1 lb. per tree. No. 2

Filler or Diluent

Sodium Aluminum Fluoride 3 lb. gal. 100 Water Liquid Blood Albumin Spreader 1/2 pt.

Peach Tree Spray

A combination of the lead arsenate and zinc-lime sprays is effective not only against chewing insects such as curculio and codling moth, but against bacteriosis. The formula is:

Zinc Sulphate	8 lb.
Hydrated Lime	8 lb.

1				
	Water Add:	100 gal.		
ı	Lend Arsennte	3 lb.		
	The sprny should be used prepared.	l as soon as		
	Prane Worm Spr	ny		
	Pyrethrum Extract Kerosene	1 qt. 6 gal.		
	Neutral Soap Water	4 lb. 94 gul,		
	Pear Tree Blight Inj U. S. Patent 2,017,			
	Pine Tar Oil Turpentine	1 oz. 16 oz.		
	Gladiolus Thrip Sp	ray		
	Manganese Arsenate (267	6 1 lb.		
	Arseme) Brown Sugar Water	66 lb. 100 gal.		
Adhesive for Hydrated Lime in Sprays				
A spray of 20 lb, calcium hydroxide and 3 lb, aluminum sulphate in 100 gal, of water will give an adherent white spray readue which is repellent to the Japanese beetle. The mixture may be of				

Lead Arsenate Substitute

value as an adherent for other spray

ingredients.

This compound is prepared by fusing 1 part diphenylamine with 2 parts sulphur at 180° C., iodine being used as catalyst. Upon recrystallizing from toluene, the light vellow crystal compound melting at 180° C., neutral, insoluble in water, slightly soluble in cold mineral oils and the usual organic solvents, is obtained. In laboratory tests, the compound is as effective as lead arsenate for codling moth larvae.

San Jose Scale Spray

1 lb. Creosote Oil Emulsion Mineral Oil Emulsion 3 lb.

Scale Insect Poison 1½ gal. 6 lb. Paraffin Oil Ferrous Sulphate Caustic Soda 20 lb, Quicklime Water 3 lb. to make 100 gal.

Holly Sprays

Use a 3% oil emulsion containing a little nicotine sulphate. This prevents scale on living trees.

Cut holly is freed from insects by dipping and soaking for 10 minutes at 24° C. in

Formalin	5) ½ gal.
Nicotine Sulphate (4%	1/5 gal.
Linseed Oil Soap	1 lb.
Water	100 gal.

Derris Spray U. S. Patent 1,934,057

Derris Extract	1-25	oz.
White Mineral Oil	40-80	oz.
Soap	5-25	oz.
Water	up to 35	oz.

The above is used diluted with water to give a mixture containing 0.06-0.25% Derris extract.

Fungi Spray U. S. Patent 2,000,843

Soft Soap	33	lb.
Cresol Soap (2% Solution)	- 11	lb.
Tobacco Extract (10%)	17	lb.
Potassium Permanganate		
(1/2 Normal)	22	lb.
Vegetable Glue	17	lb.
Alcohol	1/4-2	lb.

Lime, Sulphur and Sult Wash Formula No. 1 No. 2 No. 3 No. 4

Lime Sulphur	2 11/4	11/2	1%	2 1%	lb.
Salt	1	11/2	1%	1	lb.
Water	4	4	4	4	gal.

Boil the lime and sulphur together in a little of the water, and when combined add the rest of the water and salt. Effective as a winter application for scale.

Lime Sulphur Spray

Directions for making 50 gal, of lime sulphur spray are as follows:

Sulphur	8 lb.
Spent Carbide Residue	3 gal.
Calcium Arsenate	8 oz.

Heat about ½ of the total amount of water, adding the sulphur slowly to make a thick paste. When the water is hot, add all the carbide residue, thoroughly stirred. Mix and add another third of water and continue to cook and stir for about 45 to 60 minutes until a clear, orange-colored solution is obtained. Then add the rest of the water and the calcium arsenate. Let the mixture settle and run it through a fine sieve as it is poured into the spray tank. This should be diluted in a ratio of about six parts water to one part of the solution.

Soil Sterilization in Field and Garden Formula No. 1

The stand of such vegetables as peas, spinach and beets can usually be greatly improved by watering, immediately after planting, with a dilute solution of formaldehyde.

Formaldehyde (40%) 1 oz. Water 124 oz. Use this solution at the rate of 1 gal.

for 200 feet of row.

No. 2

Formaldehyde (40%) 15 oz. Infusorial Earth 85 oz.

When infusorial earth is used as a carrier the full strength of the formal-dehyde is maintained for a longer time than when other materials, such as charcoal or muck, are employed. Mix thoroughly, taking care to break up lumps. Use 6 oz. of this dust for each bushel of soil, or 1½ oz. per square foot of flat area. Insure that the dust is well mixed with the soil. After placing in flats, sow seed and water immediately.

Adhesives for Sulphur Dusts

Sulphur is more than twice as adhesive if applied to dry foliage; 0.25 inch of rain icmoved so nuch sulphur dust applied to dry foliage; 0.25 inch of rain icmoved so nuch sulphur dust applied to dry foliage that its effectiveness was lost. Addition of 2% of glue or gum tragacanth to dusting sulphur increased its adhesiveness 4–5 times over sulphur applied to dry foliage and twice over sulphur applied to dry foliage. When 5% of blood albumin was added to sulphur dust, its adhesiveness was increased 10 times and 5 times over that of sulphur applied alone to dry leaves and wet leaves respectively. Sulphur dust containing blood albumin remained on the leaves almost as well as did lime.

Pepper Disease Control

The use of an organic mercury dust or solution of 1 to 1000 mercuric chloride with an exposure of 5-8 minutes effectively sterilizes pepper seeds before planting. For treatment of the growing plant to control fungus diseases the use of either Bordeaux mixture or copper-lime dust is recommended. For the Bordeaux mixture, a concentration of 2-4-50 should be applied to seedbeds and 4-6-50 to more mature plants. The copper-lime dust should be mixed in the proportion of 20 lb. of dehydrated copper sulphate and 80 lb. of calcium hydroxide. These components should be mixed dry.

Dust for Control of Cucurbit Wilt
Basic Copper Chloride 1/2 oz.
Flour 5 oz.
Calcium Arsenate 1 oz.

Keep plants well covered with a light coating of dust from the time they appear through the ground until bearing stage is reached. New growth should be kept dusted. Number of applications will depend upon rate of growth and weather conditions.

Seed Disinfectant (Dustless) French Patent 770,560

Mercuric Lanolin Tale	Chloride	5	0Z. 0Z.

Lettuce Seed Sterilization

Soak 4 to 8 hours in following:
Calcium Hypochlorite 11.5 oz.
Water 1 gal.
Stir thoroughly; allow to settle; decant and use at once. Wash seeds after

Spreader for Nicotine Sprays

above treatment.

Spreaders which contain twice the amount of active ingredients and which are 4 times as effective as potassium soaps in the control of Aphis Rumics ou nasturtium leaves, are made as follows:

Formula No. 1

Water 5 g., potassium hydroxide (92%) 7.40 g., pone-tar oil (specific gravity 1.035) 44.30 g., ethylene glycol monoethyl ether 10.00 g., oleic acid 33.30 g.

No. 2

Water 5 g., potassium hydroxide 7.40 g., pine-tar oil 48.80 g., isoamyl alcohol 3.00 g., phenol (85%) 1.00 g., ethylene glycol monoethyl ether 1.50 g., and olecacid 33.30 g.

Cotton Root Rot Remedy
Apply 3% ammonium hydroxide solution.

Preventing Brown-Rot on Lemon Trees
Apply
Zinc Sulphate 40–25 lb.
Hydrated Lime 20–25 lb.
Sand 40–50 lb.

around base of trunk, piling 8 inches high and hold in place by paper collar.

Lemon Scale Control

THE PARTY OF THE COURT OF	
Yellow Sulphur	75 lb.
Gns Purification Sulphur	25 lb.
Tale	10 lb.

Grind to 200-300 mesh; use 0.4 to 1 lb. per tree at 17-day intervals. Five to seven applications are used.

Control of Cabbage Root Fly

Corrosive sublimate, applied at a strength of 1 oz. in 8 gal. of water, is the most successful means, at present known, of reducing the damage done to plants of the cubbage tribe (Brassicae) by the cubbage root fly. The treatment consists of applying to each plant about one-quarter of a pint of the solution in such a manuer as to flood the soil evenly round the base of the plants on three occasions at 10-day intervals, starting four days after setting out the plauts. Of the other methods tested, commercial naphthalene powder, about 1/4 oz., applied to the soil round the plants on three occasious at 10-day intervals commencing on the day of transplanting, possesses certain advantages, especially as regards cheapness, simplicity of application and the non-poisonous nature of the substance.

Control of Weevils in Stored Beans and Cowpens

Protection is obtained by adding 1 lb. of slaked lime per bu. of beans or cowpers and mixing thoroughly. Sodium thosalicate, used at the rate of 1 part to 1500 parts of grain, gives full protection against the grain beetle, Sitophilus granaria.

Non Poisonous Insect Exterminator Petrolatum, Laquid 1000 g. Pyrethrum Powder 200 g. Pine Needle Oil 13 g. Jumper Oil 2 g. Lavender Oil 1 g. Orange Oil 1 g.

To Kill Ants in Lawns and Gardens

Make a hole about 18 inches deep in the center of the ant hill with an old broom handle and then pour in a solution of poison made by dissolving 1 oz. of sodium or potassium cyanide in 1 gal. of water. Cover with dirt. If the soil is alkaline use one half the quantity of water and make another solution of 1 oz. of alum to 2 qt. of water and pour one-half of each in the hole.

* Deadly poison Do not allow contact with broken skin or cuts.

Beetle Powders		Endive Fly Treatmen	t
Formula No. 1		Soak roots, before planting,	
Barium Carbonate	10 oz.	20 minutes in:	40 10
Borax	20 oz.	Nicotine (50% Solution)	20 cc.
Sugar	5 oz.	Sal Soda	
No. 2	J 02.	Water	5 g. 1 l.
Sodium Fluoride	10 oz.		
Kaolin	10 oz.	Fly Dishes	
No. 3	10 02.	a. Quassia Wood	500 æ
Kieselguhr	22 oz.	Black Pepper	500 g. 50 g.
Sodium Fluoride	40 oz.	Water	2 l.
Sodium Chloride	10 oz.		
No. 4		Extract cold 4 days, then even 1 liter, filter and add:	iporate to
Powdered Borax	4 oz.	b. Sugar	100 g.
Flour	2 oz.	Color with red or green anil	
Chocolate Powder	1 oz.	Impregnate cardboard dis	hes with
No. 5		solution; dry in air.	
Powdered Borax	10 oz.		
Insect Powder	1 oz.	Killing Fly Larvae in Ces	spools
Starch	1 oz.	Add 0.15% by weight of so	
****		nide to the feeal matter.	arum cya-
Poultry Lice Powde	reg	and to the fitter matter.	
Formula No. 1		Derris Insecticide for Carawa	w Moth
Naphthalene	10 ~	Derris Root Powder	ty moun
Sulphur	10 g. 20 g.	(8% Rotenone)	1 kg.
Tobacco	40 g.	Tale	3 kg.
Talc	130 g.		
No. 2		Apply at rate of 75 kg. per two applications.	nectare in
Naphthalene	20 g.	two applications.	
Sulphur	20 g.	~	
Tobacco	40 g.	Grasshopper Poison	
Talc	120 g.	Formula No. 1	
No. 3 Naphthalene	40 æ		100 lb.
Sulphur	40 g. 20 g.	Beet Molasses	2 gal.
Tobacco	40 g.	Amyl Acetato	3 oz.
Tale	100 g.	Sodium Arsenite, Liquid	1 qt.
No. 4	6	Water No. 2	-12 gal.
Naphthalene	20 g.		100 lb.
Sulphur	20 g.	Sodium Arsenite, Liquid	1 qt.
Tobacco	40 g.	Sodium Arsenite, Powder	2 lb.
Cresol	1 g.	Water 10-	-12 gal.
Talc No. 5	119 g.		-
Naphthalene	20 g.	Groundnut (Peanut) Oil Ins	secticide
Sulphur	20 g.	This emulsion is made by n	
Sodium Fluoride	50 g.	cc. of groundnut oil with 75 c	
Talc	110 g.	acid and then pouring the mixto	
	-	with constant agitation, into a	
Dog and Poultry Flea and	Lice Killer	50 cc. of ammonia in 200 cc.	of water.
Formula No. 1		For use, this emulsion should	
Derris Powder	1/4-1 kg.	with nine times its volume of	
Water	10 1.	is stated that all insects, the w	
Shake well and rub into skin.		bodies of which are resistant to	
No. 2		aqueous liquids, are poisoned b	J tills 2/0
	1/ 1 1		
Derris Powder	1/2−1 kg.	Rat Fumigant	
Talc No. 3	10 kg.	Potassium Nitrate	30 oz.
Rotenone Solution	0.2%	Sulphur	42 oz.
2.0.010110 00.01101	/0		

Sawdust	18 oz.
Sand	6 oz.
Mix together and burn.	
Rat Bait	
Formula No. 1	
Ground Dried Bread	65 lb.
Ground Fresh Pork Fat	5 lb.
Ground Fresh Halibut or	0 10.
Haddock or Cod	20 lb.
Powdered Red Squill	10 lb.
No. 2	
Ground Dried Bread	85 lb.
Glycerin	5 lb.
Powdered Red Squill	10 lb.
No. 3	
Ground Dried Bread	37 lb.
Glycerin	3 lb.
Powdered Red Squill	10 lb.
*Fresh Bait	50 lb.
No. 4	
Ground Dried Bread 10 1	b. 10 oz.
	b. 4 oz
Zinc Phosphide	10 oz.
No. 5	
Ground Dried Bread 5 1	h
Corn Oil	10 oz.
Zinc Phosphide	10 oz.
Some Fresh Bait 6 1	b. 4 oz.
No. 6	
Ground Dried Bread 29 1	b. 6 oz.
Ground Fresh Pork Fat 2 l	
	b.
-Powdered Thallium	
Sulphate	10 oz.
No. 7	
Ground Dried Bread 18 1	
Glycerin 1 l	b. 4 oz.
Powdered Thallium	10
Sulphate	10 oz.
* Fresh bait has hamburger, or a	round sweet
potatoes (raw but canned is better applies, or ground bananas.), or ground
abbured or Product parisments.	

Rat Poison for Flour Mills Sodium Silicofluoride 70 lb. Diatomaceous Earth 30 lb.

Dust on floor, keeping away from sacks. Rats lick powder off feet and go out seeking water and thus die outside.

Rabbit Poisons

Poisoned Alfalfa. Dissolve 1 oz. of strychnine sulphate in 1 gal. of hot water and sprinkle over 10 lb. of dry alfalfa

leaves. Well-formed leaves free from dust or sticks should be used. They should be threshed thoroughly until all the moisture is absorbed. The poisoned leaves should be distributed in small handfuls, in lines a few feet apart, across portions of the field where observations show the rabbits to be feeding. Stock should be excluded.

Poisoned Green Alfulfa (summer poison) Chopped Green Alfulfa 20 lb, Strychnine (Powdered Alkuloid) 1 oz.

Saccharine 1/10 oz. Poisoned Rye Heads. In localities where alfala is not raised, rye, enimer, or wheat heads are excellent mediums for poison, and frequently surpass alfulful heaves in effectiveness, particularly in dry-land sections. Where possible, grain heads for poisoning should be cut and cured when the grain is in the dough stage, us it is more palatable and attractive to rubbits when cut at this time. Dissolve 1 oz. of strychnine sulphate in 6 gt, of hot water and sprinkle over 10 lb. of ginin heads. Mix thoroughly until all moisture is absorbed. The heads should be cut from the stem just below the last kernel and as little straw taken as possible.

Cedar Shingles.

Strychnine (Powdered
Alkaloid) 1 oz.
Saccharine 1½ oz.
Breirbonate of Soda
(Baking Soda) 1 oz.
Flour 3 tbsp.

Mix together dry, 1 oz. of powdered strychmic (akaloid), 1 oz. of baking soda, 1 teaspoonful of succharine, and 3 tablespoonfuls of flour. Add a little cold water and stir thoroughly to a smooth, creamy paste. Split the shingles and dip the tops in the paste and stick them into the ground along the rabbit trails and rinways. These baits can be easily taken up when they are no longer needed and all danger to stock is thereby climinated. In many communities this poison has proved very effective.

proved very effective.

Starch Formula (Rabbits). Dissolve 2 oz. (heaping tablespoonful) of gloss starch in a little cold water, pour into 2 to 3 quarts of boiling water, and stir until a thin starch paste is formed. Stir into the starch paste 1 oz. of strychnine (alkaloid) until a creamy paste, free from lumps, is formed. Mix the paste thoroughly over 10 lb. of grain heads until every head is coated. The heads should be cut from the stem just below

the last kernel and as little straw taken as possible. Ten pounds of alfalfa leaves or chopped alfalfa may be used in place of grain heads in alfalfa districts.

Rabbit Salt. Mix dry 1 oz. strychnine (alkaloid) with 16 oz. granulated salt. A very satisfactory method is to bore about 24 of the way through a short 2" by 4" block with 1- to 1½-inch bit and place the salt hait in this container. The blocks should be placed in or near the rabbit truils and runways. Care should be taken in placing these baits so that livestock will not obtain them.

Insect Control in Stored Rice

Fumigate with 2.5 lb. chloropicrin per 1000 cubic feet of space at temperatures above 70° F, or with carbon bisulphide at rate of 6 lb. per 1000 cubic feet.

Ground Squirrel Poison

Mix thoroughly 1 oz. strychnine alkaloid (powdered) and 1 oz. baking soda. Sift this into ¾ pint of thin, hot starch paste and stir to a creamy mass. The starch paste is made by dissolving one heaping tablespoonful of dry gloss starch in a little cold water, which is

starch in a little cold water, which is then added to ¼ pint of boiling water. Boil and stir constantly until a clear thin paste is formed.

Add ¼ pint heavy corn syrup and a tablespoonful of glycerin and stir thor-

oughly.
Add 1/8 oz. saccharine and stir thor-

oughly.
Pour this poison solution over 20 quarts of clean oats and mix thoroughly so that each grain of oats is coated. Prepare it 24 to 48 hours before using.

For mixing small quantities an ordimary galvanized wash tub is convenient. For large quantities a tight, smooth box may be used, and mixing may be done with a spade.

A teaspoonful of the poisoned oats should be placed near each ground squirrel hole on clean hard ground letting it scatter slightly as it falls. (Placed in this way it will not endanger stock). Do not put the poisoned grain on the loose dirt of the mound or into the holes. Each quart of the poisoned grain is sufficient to treat about 60 holes.

Squill Paste Preservative

A suitable preservative for the red squill paste is 1% of a hydroxybenzoic acid derivative or 1/2% of benzoic acid.

White Coal Tar Disinfectant

Cresylic Acid	50 lb.
Cresvlic Creosote	6 lb.
Sulphonated Castor Oil	5 lb.
Gelatin	3 lb.
Water	36 lb.

The sulphonated oil and gelatin are dissolved in the water and the mixed tar acids are gradually added to them with vigorous agitation in small quantities at a time. Final treatment with a colloid mill may be necessary to obtain a good dispersion.

Insecticides to Be Applied by Fumigation

German Patent 597,769

Formula No. 1	
Naphthalene	40 g.
Naphthalene, Crude	40 g.
Animal Oil	5 cc.
Cresol, Crude	5 cc.
Ceresin	10 g.
No. 2	6 .
Naphthalene	15 g.
Naphthalene, Crude	25 g.
Animal Oil	20 cc.
Cresol, Crude	30 cc.
Ceresin	10 g.
No. 3	
Naphthalene	10 g.
Naphthalene, Crudo	10 g.
Animal Oil	25 cc.
Cresol, Crude	25 cc.
Ceresin	30 g.

Method: Melt naphthalene and ceresin, and add cresol and oil at low temperature. Application on fumigation-pans.

Warehouse Fumigant

Chloropicrin (nitrochloroform, CCl₃·NO₂) is a colorless heavy liquid which is becoming prominent as a funigant. Due to its highly lachrymatory nature as well as its highly toxic effect on insects, their larvae and eggs, it makes possible effective fumigation without the high possibility of accidental death to operators attendant on the use of hydrogen cyanide.

The following are the recommended quantities in pounds per 1000 cu. ft. of volume to be funigated using 24-hour exposure and a temperature of 70-80° F. Higher temperatures reduce the exposure time while lower ones increase it. The liquid is vaporized by spraying or evaporation from very shallow pans or soaked

cloth.

		1	
Confectionery Industry.	Lb. per	Calcium Hypophosphite Sodium Hypophosphite	8 g. 12 g.
	1000 cu. ft.	l Water, Distillate	71 g.
Candy Nut s	1 1	No. 3	
	•	35% Oil	
Dairy Industry.	1/	Carragheen Moss	18 g.
Eggs and Cheese	1/4	Distilled Water	400 g.
Milling Industry.		Sodium Formate	5 g.
Macaroni Vaults	1-11/4	Cod Liver Oil	350 g.
Macaroni, Cased	11/2-2	Syrup, White Distilled Writer	100 g. 117 g.
Space Fumigation Flour Mills	1	*Spice Oil Mixture	10 g.
		•	10 6.
General Returned Bags	1 114	No. 4	
Sacked Flour	11/2	30% Oil	10
Fly and worm control.	- /4	Gum Arabic Gum Tragacanth	12 g. 16 g.
(Exposure over the week en	nd) 1/4	Glycerm (28° Bé.)	140 g.
Rice Bags and Vaults	114	Distilled Water	430 g.
Box Cars (Adults)	4-5	Moldex or Other Good Pre-	
Box Cars Complete	7-8	servative	1 g.
Stored Products.	1	Cod Laver Oil	300 g.
Warehouse Space	1	*Spice Oil Mixture Calcium Hypophosphite	10 g. 8 g.
Sacked Goods and Vaults	114	Sodium Hypophosphite	12 g.
Grain Bins		Distilled Water	71 g.
(With grain moving at 100)	No. 5	.,
bu. per hr.)	2	f Gum Arabic	15 g.
Contaminated Bins	3	Gum Tragacanth	8 g.
l'obacco.	1	Glycerin (28° Bé.)	50 g.
Vaults	2	Distilled Water	456 g.
Warehouses	11/4	Sodium Formate	5 g.
Furniture.		Iodine Chloroform	3 g. 3 g.
Furniture	11/4	*Spice Oil Mixture	3 g.
Household.		Cod Laver Oil	447 g.
Bedbugs, Clothes Moths,		No. 6	
Roaches	1	(Gum Arabic	10 g.
Buffalo Moth	11/4	Gum Tragacanth	10 g.
Rodents	1/4	Glycerin (28' B6.)	200 g.
Cod Liver Emulsion for A	nimals	Water, Distilled Potassium Todide	366 g. 3 g.
Formula No. 1		Moldex or Other Good Pre-	о в.
50% Oil		servative	1 g.
Carragheen Moss	12 g.	Cod Liver Oil	400 g.
Distilled Water	300 g.	*Spice Oil Mixture	10 g.
Moldex or Other Good Pre-		No. 7	
servative	1 g.	Carragheen Moss	19 g.
Cod Liver Oil	500 g.	Glycerin (28° Bé.)	100 g.
Syrup, White Distilled Water	86 g. 91 g.	Distilled Water	519 g.
*Spice Oil Mixture	10 g.	Potassium Iodide Moldex or Other Good Pre-	1 g.
•		servative	1 g.
No. 2		Cod Liver Oil	350 g.
40% Oil	10 ~	*Spice Oil Mixture	10 g.
Gum Arabic	12 g. 12 g.	No. 8	
Gum Tragacanth Glycerin (28° Bé.)	130 g.	Gum Arabic	12 g.
Water, Distilled	340 g.	Gum Tragacanth	16 g.
Sodium Salicylate	5 g.	Glycerin (28° Bé.)	130 g.
Cod Liver Oil	400 g.	Distilled Water	426 g.
*Spice Oil Mixture	10 g.	Sodium Salicylate	5 g.

Iodine	1 g.
Alcohol. Absolute	10 g.
Cod Liver Oil	300 g.
Spice Oil Mixture	10 g.
Calcium Hypophosphite	8 g.
Sodium Hypophosphite	12 g.
Uistilled Water	70 g.
Spice Oil Mixtures for Above	Emulsions

Formula No. 1

Vermonth Oil	5	CC.
Coriander Oil	2	cc.
Galanga Oil	1	cc.
Gentian Oil	1	cc.
Culanus Oil	0.5	ee.
Peppermint Oil	0.5	ec.

Τ,	O.	4	

Fennel Oil	5 cc.
Calamus Oil	3 cc.
Peppermint Oil	2 cc.

No. 3

Fennel Oil	6 cc.
Calirmus Oil	4 cc.

The above emulsions are made up best in enameled kettles with high speed mixers.

Gnm Solutions: Wash gnm arabic with water at 40° C, then put into cold water and warm to solution. Gnm tragacanth or enringheen moss are first wet with glycerin and put into cold water. Soak 12 hours. Prepare gnms separately and when ready, mix as indicated, warm up to 90° C, (add Iodide) then add preservative.

Stir in Cod Liver Oil in small portions. Then add Spice Oil Mixture, with stirring, Syrup and Hypophosphites are dissolved in hot water as indicated. Stir into cumbion hot. Iodine is prepared by solution in Alcohol or Chloroform and a little of the Cod Liver Oil, then is added to the gnm (aqueous) solutions and emulsified.

When ready, stir vigorously for 1/2 hour, or put through a homogenizer.

Cod Liver Oil Emulsion for Animals

Formula No. 1

rormula No. 1	
(Gum Arabic	100 g.
	100-120 g.
l Glycerin	1200 g.
 b. Cod Liver Oil, Crude 	3700 g.
Calcium Hypophosphito	50 g.
c. Sodinm Hypophosphite	50 g.
Water	4000 g.

Grind a until smooth, add b in small portions, homogenizing every time. To this add c in an emulsifying machine.

As spice, add 1% of the following mixture of:

Vermouth Oil	10 cc.
Coriander Oil	4 cc.
Galanga Oıl	2 cc.
Gentian Oil	2 cc.
Calamus Oil	1 cc.
Peppermint Oil	1 cc.

No. 2

1.0. m			
a. { Iceland Moss Water (2 portions)		10	g.
Water (2 portions)	to	600	g.
			Extract
Gum Tragacanth		6	g.
b. Gum Tragacanth Gum Arabic Cod Liver Oil		6	g.
Cod Liver Oil		400	g.
c. Fennel Oil		5	drops
Calamus Oil		5	drops

Boil a two times (two portions of water) to 600 g. nnited extract. Grind b nntil homogeneous and transfer into a dry bottle; add c, then a in two portions, shaking thoroughly and vigorously.

No. 3

110. 0	
Carragheen Moss	10 g.
a. Carragheen Moss Water	350 g.
b. Cod Liver Oil	500 g.
(White Syrup	100 g.
c. {White Syrup c. {Malt Extract	20 g.
Water	120 g.

Soak a for 12 hours, boil then about 10-15 mm., filter through cloth. Add b, while stirring, to this hot solution, then stir in c, and add as preservative

Sodium Salicylate 0.3-0.5 g.

No. 4

	110. 1	
a.	Gum Tragacanth	5 g.
	Gunt Arabic	8 g.
	Water	250 g.
b.	Calcium Chloride	50 g.
	Water	57 g.
c.	Lame Water	230 g.
	Cod Liver Od	400 or

Souk a for $1\frac{1}{2}-2$ days, add b, then a, mix well, percelate (lumps remaining on the cloth are ground with water and pour again through the filter). Mix the whole well in an emulsifying machine with d for hours; d is added in 8 portions. The a, b, c is treated alone before.

Skin Abrasion Lotion (For Dogs)

Dissolve 1 part of castile soap in 9 parts of water. Wash dog thoroughly with this solution; and then apply with cotton to the affected parts 5% tincture of iodine.

Mois	ture	Ecze	ma	Lotion	for	Dogs
This	lotio	n is	exe	cellent	for	bathing

moist eczema spots on dogs.

 Tannie Acid
 5 oz.

 Salicylie Acid
 5 oz.

 Alcohol (50%)
 90 oz.

Before using this preparation the spots should be thoroughly washed with castile soan.

Dog Eczema Powder

Senega Root Powder 90 oz.
Sodium Sulphite 10 oz.
Rub into skin with water and finally wash off.

Liquid Soap for Dogs and Other Animals
Palm Kernel Oil 1200 g
Olein 200 g
Caustic Potash (50%) about 736 g
Glyeerin 600 g.
Softened or Distilled Water 6800 g.
Carbolic Acid (Phenol),
Crude 400 g.
Perfume Oil (e.g., Euca-

Dog Deterren

lyptus)

50 g.

Dog Deterrent		
Naphthalene Flakes	4	οZ
Paraffin Wax	1/4	OZ.
Gasoline	1 - 2	
Rosin	1/4	07.

Stir until dissolved; spray on base of tree trunks or shrubs with an insect spray gun-

Dog Nuisance Preventer

To prevent dogs from stanning trees and shrubs, spray the base of the latter with a solution of $\frac{1}{1}$ oz. nicotine sulphate per gal, water.

Dog Worm Remedy

roimun 100 1	
Aloes	45 gr.
Soap	45 gr.
Oleoresin of Male Fern	30 gr.
Mix and make into 2 pills.	

Administer both pills in the morning, the animal to remain fasting for some time.

No. 2

Areca nut, freshly ground, is considered an excellent remedy for worms in dogs. About one dram made into a pill is the dose for an ordinary sized dog. This should be given at night followed by a dose of castor oil in the morning.

Animal Eve Washes

One of the best eye washes for irrigation and cleansing of the eye and for puralent discharges and conjunctivitis is as follows:

Sodium Bicarbonate	15 gr.
Borax	15 gr.
Sodium Chloride	15 gr.
Glycerin	1 dr
Distilled Water	8 oz.

Animal Ear Preparation

Formula No. 1

	I Official 240, 2		
Gentian	Violet	5	υŻ
Acctone		5	07.
Alcohol		45	07.
Water		45	07.

Take small amount in an eye dropper and place deep into the ear and remove excess so as not to soil the outside.

	No. 2	
Phenol		3 oz.
Glycerin		97 oz.

Add boric acid powder until the glycerin will not absorb any more. Let stand over night and strain.

Place one half eve dropperful in eye and remove the excess.

Dog Mange Treatment

Formula No. 1

Kerosene	32 oz.
Creolin	6 oz.
Oil of Tar	6 oz,
Sulphur	1 lb.
Raw Linseed Oil	to make 1 gal.
Rub into skin eve	ry other day. It
gves gratifying resu	lts.

No. 2

Another good only skin	mixture is:
Gum Camphor	1 lb.
Alcohol	1 pt.
Turpentine	1 qt.
Kerosene	2 qt.
Cotton Seed Oil	6 qt.
Sulphur (Howers)	9 oz.

Note: First dissolve the camphor in the alcohol. Rub on the skin every third day.

Dog Mouth Wash

Tincture Iron			1	OZ.
Potassium Chlorate			2	OZ.
Glycerin			4	OZ.
Water	to	make	1	gal.

Aphrodisiac for Cattle and Horses

The usual doses of yohimbine hydrochloride as an aphrodisiac in veterinary practice are: Stallons, 1 gr.; bulls, 1¼ gr.; cows and mares, 1½ gr. It should be administered in the food or drinking water three times a day.

Cow Abortion Flush

Common Salt 1 lb. Potable Water 95 lb.

Remove aborting cow from herd. Before returning to herd flush daily with above solution.

Bloody Milk Mixture

Glauber's Salts 1 lb. Water 4 lb.

Give the above dosage to cow producing bloody milk. Find and remove the cause; it may be udder injury, improper feeding, or overfeeding. Certain bacteria inpart a red color to milk; this is uncommon.

Cow Boil Wash

Carbolic Acid Solution (3%)

Syringe out cavity with above solution after lancing and removing contents,

Chapped Teats Solution

Boric Acid Crystals 1 lb. Water 15 lb.

Bathe teats twice daily with above and dry; then rub teats with vaseline.

Cow Pox Solution

Apply a 4% solution of potassium permanganate after cleaning udder and teats.

Calf Scours Remedy

Salol 1 lb. Subnitrate of Bismuth 2 lb.

First give the calf with simple scours 1½ oz. of castor oil in ½ pt. of warm milk. After a few hours give a teaspoonful of the above. Repeat this dosage three times daily.

Impaction in Cattle Treatment

Glauber's Salts 1½ lb. Water 7 lb.

Administer 2 oz. of aromatic spirits of ammonia at once. Two hours later give the above formula.

Egg Preserving Solution

Sodium Silicate 1 fl. oz. Water 25 fl. oz.

Defeathering Poultry U. S. Patent 2,017,648

Burgundy Pitch 15 lb. Montan Wax 5 lb. Paraffin Wax 10 lb.

The ingredients are melted, thoroughly mixed, and applied to the careass, preferably after the bird has been scalled and the bulk of the feathers that can be removed hastily have been removed by hand.

After application, the defeathering compound is permitted to solidify by cooling and is then removed, taking with it epidermal excrescences such as feathers, down, pinfeathers and the like.

Prevention of Skin Tearing when Plucking Feathers with Adhesives

Spray the skin with an oil emulsion.

Bird Gravel

Fine River Sand	97.5 g.
Cuttlefish Bone, Powder	2 g.
Pyrethrum Flowers	0.5 g.

Ration No. 1 (With Milk)

For those who wish to use home-grown grains.

18 lb.
18 lb.
18 lb.
18 lb.
10 lb.
10 lb.
5 lb.
2 lb.
1 lb.

Ration No. 2 (With Milk)

Using wheat by products.	
Ground Corn	20 lb.
Bran	20 lb.
Flour Middlings	12 lb.
Ground Oats	20 lb.
Meat Scrap	10 lb.
Dried Milk	10 lb.
Alfalfa Meal	5 lb.
Steamed Bone Meal	2 lb.
Salt	1 lb.

This is a ration for those who wish to use barley in the laying ration. Barley

is not as palatable as corn when fed whole in the scratch grain but is a valuable ingredient of a laying mash. However, it should be remembered that this grain is low in vitamin "A" when compared with corn and that sufficient alfalfa meal should be present to take caro of this deficiency.

Ration No. 3 (With Milk)

(
Ground Barley	20 lb.
Bran	20 lb.
Flour Middlings	10 lb.
Ground Oats	20 1Ь.
Meat Scrap	10 lb.
Dried Milk	10 lb.
Alfalfa Meal	7 lb.
Steamed Bone Meal	2 lb.
Salt	1 lb.

Ration No. 4 (Without Milk)

(1116000	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
Ground Corn		20	lb.
Bran			lb.
Flour Middlings			lb.
Ground Oats			lb.
Meat Scrap			lb.
Alfalfa Meal			lb.
Salt		1	lb.

Ration No. 5

Many farmers and poultrymen wish to feed a surplus of liquid milk (either skim or buttermilk) to the laying flock. This is a successful practice and the following ration is designed to be fed when liquid milk is given as the only drink. In omitting the water for drinking purposes no fear need be felt as liquid milk is about 90% water. If water is given to the flock in addition to the liquid milk, the meat scrap content should be in creased to obtain best results. It should also be remembered that the practice of feeding liquid milk for one or two days and then missing a day is a bad one and a satisfactory production cannot be expected over a long period of time.

Ground Corn	20 lb.
Ground Oats	21 lb.
Ground Barley	21 lb.
Ground Wheat	20 lb.
Meat Scrap	10 lb.
Alfalfa Meal	5 lb.
Steamed Bone Meal	2 lb.
Salt	1 lb.

The above ration may also be used when condensed buttermilk is fed.

Ration No. 6

The following ration is one which has been fed at the Poultry Experiment Station to 1200 laying hens during the past year and the egg production and hatch-ability obtained have been satisfactory.

Ground Barley	28 16.
Bran	20 1ь.
Ground Oatmeal	11 lb.
Flour Middlings	10 lb.
Ment Scrap	10 lb.
Dried Milk	10 lb.
Alfalfa Leaf Meal	8 lb.
Stenned Bone Meal	2 lb.
Salt	1 lb.

Either dired buttermilk or dried skim milk may be used in unking up these laying rations. If one is feeding for eggs alone, any of the foregoing nations will give good results. If good hatchally is desired, rations No. 1, No. 2, No. 3 and No. 6 are recommended.

A satisfactory scratch grain consists of equal parts, by weight, of corn and wheat.

It is important that pullets especially obtain enough scratch grain to keep them in good growing condition. They are inder the double strain of egy production and growth. Do not obtain full and winter eggs from your pullets at the expense of growth as this leads to monit.

Oyster shell should be available to the laying flock at all times.

All-Mash Ration for Laying Hens

Yellow Corn, Coarsely		
Ground	35	lb.
Wheat, Coarsely Ground or		
Shorts	30	Ilı.
Oats, Finely Ground	20	lb.
Wheat Bran, Course	7	lb.
Ment Scrap, Medium		
(50-55% Protein)	10	lb.
Dried Skim Milk or Butter-		
nulk	3	lb.
Alfalfa Meal or Leaf Meal	5	lb.
Salt	0.5	lb.

In addition, for confined layers, use ½ to 1 pt., or amount suggested by manufacturers, of potent cod liver oil or sardine oil to each 100 pounds of mash. In case it is preferred that grain amash be fed separately, the following

formulas may be used:

Mash Ration

Coarsely Ground Yellow Corn	20	lb.
Wheat, Coarsely Ground or		
Shorts	20	lb.
Onts, Finely Ground	20	lb.
Wheat Bran, Coarse	9	lb.
Meat Scrap, Medium		
(50-55% Protem)	20	lb.
Dried Skim Milk or Butter-		
milk	- 5	114

100	HIMITON	II FORMODAICI	
Alfalfa Meal	5 lb.	Dried Skim Milk or Butter-	
Salt	1 lb.	nulk	5 lb.
Cod Liver Oil	1 lb.	Alfalfa Meal or Leaf Meal	
	1 10.	Cod Liver Oil	1 lb.
Grain Ration		our Eiver on	
Whole Wheat	2 lb.		
Whole or Cracked Corn	2 lb.	Ration for Fattening Chi	ckons
Whole Oats or Barley	1 lb.		скень
or		Formula No. 1	
Whole Wheat Whole or Cracked Corn	. 1 4 -	Finely Ground Corn	12 lb.
Whole or Cracked Corn (equa	ai parts	Wheat Bran	4 lb.
•		Wheat Middlings	4 lb.
		Meat Scrap	1 lb.
Egg-Laying Rations		No. 2	
****		Finely Ground Oats	15 lb.
Mixture No. 1		Finely Ground Corn	15 lb.
Mash:		Low Grade Flour	2 lb.
	16 lb.	Bran	1 lb.
Meat Scrap	6 lb.	To fatten chickens, feed of	ne of the
Bran	1 lb.	above mixtures 3 times da	
Middlings	1 lb.	should be made soft with but	
Sciatch Mixture:		skini milk.	
Cracked Corn	1 lb.	***************************************	
Wheat	1 lb.		
No. 2		Breeding Flock Ratio	n
Mash:		Mash:	
Barley Meal	2 lb.	Bran	1 lb.
Bran	1 lb.	Middlings	1 lb. 3 lb.
Middlings	1 lb.	Corn Meal Meat Scrap	3 lb.
Fish Scrap	1 lb.	Meat Scrap	11/2 lb.
Scratch Mixture:		Ground Oats Rolled Oats	1 lb.
Cracked Corn	1 lb.		1 lb.
Wheat	1 lb.	Linseed Meal	⅓ lb.
377141 41 - 1 - 1 - 1		Scratch Mixture:	
With the above mixtures sup		Cracked Corn	1 lb.
green feed. Feed scratch mixt		Wheat	1 lb.
daily and sparingly. Feed sers	l loto m	Keep breeding stock outd	oors every
the afternoon. Mash may be	fod dry	good day throughout the yea	r. Suppiy
or wet.	red dry	abundance of green feed. Fo	eed scratch
		feed in deep litter to make her	
		Fertile eggs can be produce	d by not
Chick Feed		forcing the hens with food, an	
Yellow Corn Meal (Ground		ing vigorous males also well fe	1.
Coarsely)	360 lb.		
Bran	200 lb.	Poultry Appetite Stimu	lant
Ground Ontmenl	200 lb. 200 lb.	Pulverized Gentian	1 lb
Skim Milk Powder	100 lb.	Pulverized Ginger	1/4 lb.
Meat Serap	50 lb.	Pulverized Saltpeter	1/4 lb
Alfalfa Long Monl		1	.7
	50 lb.	I Pulverized from Sulphate	- 1/6 1b.
Steamed Bonemeal	50 lb. 20 lb.	Pulverized from Sulphate Pulverized Nux Vomica	⅓, lb. 1/4 lb
Steamed Bonemeal		Pulverized Gentian Pulverized Ginger Pulverized Saltpeter Pulverized Iron Sulphate Pulverized Nux Vomica Add Log of the proposition	1/2 lb.
	20 lb.	Pulverized from Sulphate Pulverized Nux Vomica Add 1 oz. of the preparation 5 lb. of mash.	½ lb. ¼ lb. on to each

Chick Starter Feed		
Yellow Corn, Coarsely Ground	50	lb.
Coarsely Ground Wheat or		
Middlings	20	lb.
Wheat Bran	10	lb.
Meat Scrap (50-55% Pro-		
tein)	10	lb.

Poultry Coccidiosis Feed Skim Milk or Butter-

Dry Skim Milk or Butter-		
milk	40	lb.
Wheat Bran	10	lb.
Yellow Corn Meal	30	lb.
Ground Barley	20	lb.
Ferrous Sulphate	1/4	lb.

Powder for Hens to Increase Egg Production

Formula	No.	No 2	No.
Dicalcium Phosphate,	g.	g.	g.
Precipitated	72	70	
Calcium Carbonate			60
Ferrous Sulphate, Pow-			ĺ
der	12	10	
Ferrous Oxide, Powder.		-	10
Black Pepper, Ground	6		5
Ginger Root, Powder		20	10
Gentian Root, Powder	10		
Stinging Nettle Seed			15

Harrison Test Cow Feed

This is the formula recommended by Cornell University for test cows—It can be successfully used with second cutting alfalfa or second cutting timothy and clover.

Formula No. 1

roimune 110. 1	
Distillers Grain (9% Fat)	300 lb.
Wheat Bran	400 lb.
Hominy or Corn Meal	400 lb.
Ground Oats	370 lb.
Coconut Oil Meal	300 lb.
Linseed Oil Meal	200 lb.
Steam Bone Meal	20 lb.
Salt	10 lb.

(18% protein feed.) No. 2

Soybean Feed

This can be successfully fed with good hay.

Ground Oats Ground Soybeans	900 100	
--------------------------------	------------	--

Fattening Powder for Pigs

Formula	No 1	No.	No.
	g.	g.	g.
Salt	11	20	10
(8b ₂ S ₃ crude)	10	10	
Sulphur Flowers	11	10	
Glauber's Salt, Crystal-	11	20	10
Sodium Bicarbonate	21	217	
Trigonella Seed	16	10	20
Linseed Meal	20		
Fennel, Pulverized		10	
Gentian Root Powder		10	13
Juniper Berries, Dry, Pow-			
der		10	20
Calamus, Powder	-		14

Milk-Increasing Powder for Cows Formula No. 1

Calcium Carbonate	50 g.
Caraway Seed	30 g.
Calamus, Powder	20 g.
No. 2	
Dicalcum Carbonate, Pre-	
cipitated	40 g.
Caraway Seed	20 g.
Calamus, Powder	20 g.

Goat Feeds

1. Ground Feed for Bucks

Trigonella Seed

100 10.
100 lb.
50 lb.
25 lb.

Feed at rate of 1½ lb, per buck duily; mercase to 2 lb during breeding season, belode 3 lb, of alfalfa or clover hay, and a pound of turops with the ration of ground feed.

2. Vorbies Grain Mixtures for Does

I.	
Rolled Barley	100 lb.
Wheat Bran	100 lb.
Dried Beet Pulp	100 lb.
Coconut Oil	100 lb.

Feed 1 to 2 lb, per doc daily along with hay and mangels, $\frac{1}{2}$

600 lb.
100 lb.
100 lb.
200 lb.

Feed 1 to 2 lb. per doe daily along with hay and turnips.

100 11

III.

Wheat Bran	100 lb.
Oats	100 lb.
Coconut Meal	100 lb.
Feed 1 to 2 lb, per do	e per day alon

Feed 1 to 2 lb. per doe per day along with hay and turnips or silage.

17.	
Dried Beet Pulp	300 lb.
Rolled Barley	100 lb.
Wheat Bran	100 lb.

Feed 1 to 2 lb. per doc per day along with bay and mangels.

California Kid Feed Formula

Rolled Barley	100 lb.
Ground Oats	100 lb.

Peed 1/4 to 1/2 lb. daily per kid after two weeks of age. Allow animals to eat hay, and give milk.

Feeding Lime for Animals

No. 1	No. 2	No.
g.	g.	g.
65	70	80
10	-	5
6	9	3
4	4	3 3
	4	3
4	3	3 3
7	9	3
	g. 65 10 6 4 4	65 70 10 — 9 4 4 4 4

Pasture Seed Mixture

Formula No. 1

Timothy	40 lb.
Alaike Clover	20 lb.
Kentucky Bluegrass	20 lb.
White Clover	20 lb.
Orchard Grass	20 lb.
Redtop	20 lb.
Meadow Fescue	20 lb.

The above formula is used for seeding pastures not to be layed. Use 16 lb. of formula per acre.

No. 2

For Wet and Unproductive	Land
Alsike Clover	20 lb.
Canada Bluegrass	40 lb.
White Clover	20 lb.
Orchard Grass	40 lb.
Redtop	40 10.
Use 16 lb. per aere.	

No. 3	
Timothy	80 lb.
Red Clover	20 lb.
Alsike Clover	20 lb.
Kentucky Bluegrass	20 lb.
White Clover	20 lb.
Redtop	20 lb.
Orchard Grass	20 lb.

Use 20 lb. per acre. For a year or two the field should be hayed. After that when the plants are firmly established it should be pastured.

Garden Fertilizer

Nitrate of Soda	135 lb.
Sulphate of Ammonia	200 lb.
Animal Tankage	250 lb.
Superphosphate	1000 lb.
Muriate of Potash	200 lb.
Filler	215 lb.

Fertilizer

Formula No. 1 French Patent 779.281

richen raceas	
Calcium Phosphate	75 lb.
Gypsum	20 lb.
Sulphur	5 lb.

No. 2

U. S. Patent 1,931,296

Roast following mixtures for 30 min-

ites at 31	9-429 r.		
a. Rock	Phosphate	40	lb.
Lime		10	lb.
Salt		21/2	
b. Coal		35	lb.
Salt		21/2	lb.
Grind a	bove with		
Ammon	ium Sulphate	10	lb.

No. 3

British Patent 410,487

Moist Sewage Sludge	50 lb.
Chalk	15 lb. 5 lb.
Slaked Lime	
Dust, Refuse, Etc.	30 lb.

No. 4

U. S. Patent 2,019,713

Ammonium nitrate and ammoniated triple superphosphate in the proportions of about 45 to 60 parts of ammonium nitrate to 55 to 40 parts of ammoniated triple superphosphate.

Plant Food

X 1(0.10 = 0.00	
Trisodium Phosphate	2 oz.
Potassium Sulphate	2 oz.
Sadam Nitrate	3 oz.

Grind together and mix well. Only about a half gram of the above mixture should be used per plant every month or two. Caution: Using too much of any plant food is dangerous.

House Plant Food

Potassium Nitrate (Salt-

peter) 3 oz.
Tribasic Sodium Phosphate 2 oz.

Mix, and dissolve about one tablespoon to the gallon. Of this solution, use one gill for each average size plant, once every two weeks.

Alkali Farm Land Treatment

"Alkah" spots on western farm land are usually due to the presence of sodium clay. Finely pulverized gypsum (calcium sulphate), thoroughly worked into the soil over a period of a year, will usually prove an effective remedy.

Detecting Treated Grains

Limed grain may be easily detected by the red color developed when it is dropped into a dilute solution of phenol-phthalein.

Sulphur bleached grain may be de-tected by the dark color developed when

it is dropped into a dilute solution of lead acetate or lead nitrate.

Delinting Cotton Seed

Seed having a moisture content of 7 to 10% is treated with hydrochloric acid (2% on weight of seed) up to 60° E for 7 minutes. Treatment at 20° E requires 15 to 30 minutes.

FOOD PRODUCTS, BEVERAGES, FLAVORS

Ice Cream

Formulas are presented for seven series of ice-cream mixes containing 20 to 50% cream, showing the proportions of whole, skimmed, condensed, or dried milk that must be mixed in various combinations to produce the desired percentage of solids in the ice cream. These formulas show the ratios of milk fat to scrum solids which are commonly used for different types of ice cream.

In Tables 1 to 5, the formulas contain the following dairy products, with 15% sugar added to the mixtures and 0.3% gelatin:

- 1) Cream, skim milk, and whole milk. (2) Cream, unsweetened condensed skim milk, and either skim milk or
- whole milk.
- (3) Cream, dry skim milk, and either skim milk or whole milk.
- (4) Cream, sweetened condensed whole milk and skim milk.
- (5) Cream mixed with 50% butter, to which mixture is added either dry or unsweetened condensed skim milk, making ice cream containing about 6 to 11% butter.

Tables 6 and 7 show combinations of dairy products without the addition of sugar which are suitable for basic mixes in milk plants for shipment to ice cream manufacturers. In each case, 15 lb. of sugar should be added to each 85 lb. of the unsweetened mix, in order to make a palatable commercial product. The dairy products used in these two tables are:

- (6) Cream, unsweetened condensed skim milk, and either whole or skim milk.
- (7) Cream, dried skim milk, and either whole or skim milk.

For each of these formulas there are given:

- (A) Percentage of solid constituents desired in the ice cream to produce ice cream containing 10 to 18% fat, from 20 to 50% ice cream;
- (B) Groups of ingredients which may be used in making comparable ice creams of the same solids content;
- (C) The percentages by weight of each of the different milk products required to give a mixture of the desired solids content.

The quantity of each ingredient needed for different size batches of the various mixtures can easily be determined by multiplying the quantity of the total mixture desired in pounds, by the percentages given in the tables.

The flavor and texture of ice cream will vary according to the proportion of milk solids, sugar, gelatin, and flavoring materials present, and the quality of the ingucdients used. Ice-cream makers should therefore be careful to select ingredients and ice-cream formula of character and type best suited to their trade, and should check the accuracy of their figures in proportioning each mixture.

Ice Cream

Formula No. 1

100 lb. Mix--8% Fat-Cream, Whole Milk and Skim Milk Powder, 12%

Sugar.	
Sugar	12 lb.
Gelatin	0.5 lb.
Skim Milk Powder	6 lb.
Cream (30%)	18.5 lb.
Milk (1%)	63 lb.
No 9	

8% Fat-Cream Skim Milk, Skim Milk Powder, 12% Sugar.

Sugar	12 lb.
Gelatin	0.5 lb.
Skim Milk Powder	6 lb.
Cream (30%)	26.7 lb.
Skim Milk	55.8 lb.
No. 3	

8% Fat-Sweet Butter, Whole Milk,

Skim Milk Powder, 12%	sugar.
Sugar	12 lb.
Gelatin	0.5 lb.
Skim Milk Powder	6 lb.
Butter (84%)	6 lb.
Milk (4%)	75.5 lb.
No. 4	

8% Fat—Sweet Butter, Skim Milk, Skim Milk Powder, 12% Sugar.

Mille I Uwuci, 1270	Bugai.
Sugar	12 lb.
Gelatin	0.5 lb.
Skim Milk Powder	6 lb.
Butter (84%)	9.6 lb.
Skim Milk	72 lb.

Table 1.—Amounts of Cream of Different Fat Content and Either Skim Milk or Whole Milk Necessary for Making Different Types of Ice Cream

A. Solids	Types of Ice Cream					
	а	ь	o	d	e	
	Per Cent	Per Cent	Per Cent	Per Cent	Per Cent	
Fat	14	16	16	17	18	
Serum Solids	6.39	6.30	6.21	6.12	6.03	
Sugar	15.00	15.00	15.00	15.00	15.00	
(ielatın	0.3	0.3	0.3	0.3	0.3	
Total Solids	35.69	36,60	37.51	38.42	39.33	
B. Ingredients Per cent C	r cent C. Percentage of Ingredient by Weight fat Ice Cream Mixture				ach Type of	
	a	b	c ·	d	c	
1 Creum 50	28 00	30,00	32.00	34.00	36.00	
Skim Milk 0	57.00	55 00	53.00	51.00	49.00	
2 Cream 40	35.00	37.50	10.00	42.50	45.00	
Skim Milk 0	50 00	47,50	45.00	42.50	40.00	
3 Cream 30	46.75	50.00	53,50	56.75	60 00	
Skim Milk 0	38 25	35.00	31.50	28 25	25.00	
4 Cream 20	70.00	75.00	50,00	85.00	****	
Skim Milk 0	15.00	10.00	5,00			
DIG. 1.2.1.	23.00	25 25	27.50	29.75	32.00	
5 Cream 50 Whole Milk 4	62,00	59.75	57.50	55 25	53.00	
	29.5	32 25	35 00	37.75	40.50	
6 Cream 40 Whole Milk 4	55 5	52.75	50,00	17.25	41.50	
Whole Mills III	40.75	11.75	48 50	52 50	56,50	
7 Cream 30 Whole Milk 4			36,50	32.50	28.50	
	44 25	40 25			20.00	
8 Cream 20	66 75	72.50	78.75	85 00		
Whole Milk 4	18.25	12.50	6 25			
Add to each above com-						
bination:	15 00	15,00	15.00	15 00	15.00	
Sugar	0,3	0.3	0.2	0.3	0.3	
Gelatin		100.00	100 00	100.00	100.00	
Total	100.00	100,00	100 00	100.00	700.00	

Note: Ice creams made from these formulas whip and freeze slowly, and are likely to develop a buttery consistency, especially if the temperature is not kept fairly constant during storage in the hardening room or cabinet. The use of homogenized cream or mix will prevent undesirable fat clumping in freezing. Aging of the inixes for 24 hours at 40-50° F, before freezing will improve the texture. The cream flavor will be especially noticeable in the high-fat ice creams, hence care should be taken to use only high-grade cream. Melting will be accompanied by leaking of a milky serum from the ice and whipped cream structure of these ice creams, which keep their original form to a considerable extent instead of melting in a homogeneous mass. This is a natural characteristic of straight-cream ice creams, and does not constitute a defect.

No. 5 8% Fat—Sweet Butter, Sk der, Water, 12% S	im Milk Pow-	No. 6 8% Fat—Cream, Whole I Powder, 14%	Milk, Skim Mill Bugar.	k
Sugar	12 lb.	Sugar	14 lb.	
Gelatin	0.5 lb.	Gelatin	0.5 lb.	
Butter (84%)	9.6 lb.	Skim Milk Powder	6 lb.	
Skim Milk Powder	13.2 lb.	Cream (25%)	23 lb.	
Water	64.7 lb.	Milk (4%)	56.5 lb.	

TABLE 2.—Amounts of Cream of Different Fat Content; Unsweetened Condensed Skim Milk, and Fresh Skim Milk or Whole Milk Necessary for Making Different Types of Ice Cream

A. Solids	Types of Ice Cream						
	a	b	c	d	e		
	Per Cent	Per Cent	Per Cent	Per Cent	Per Cent		
Fat	10 11 15	12 10 15	14 9 15	16 8 15	18 7 15		
Gelatin	$0.3 \\ 36.3$	$\frac{0.3}{37.3}$	$\frac{0.3}{38.3}$	$\frac{0.3}{39.3}$	0.3 40.3		
		of Ingre		eight in I	Each Type of		
	a	b	c	đ	e		
1 Cream 50 Skim Milk 0	20.00 41.00	$24.00 \\ 42.50$	$28.00 \\ 42.50$	$\frac{32.00}{43.00}$	36.00 43.75		
2 Cream 40 Skim Milk 0	$25.00 \\ 36.00$	$\frac{3000}{36.50}$	35.00 35.50	40.00 35.00	45.00 34.75		
3 Cream 30 Skim Milk 0	33.30 27.70	$\frac{40.00}{26.50}$	$\frac{46.70}{29.80}$	$53.40 \\ 21.60$	60.00 19.75		
4 Cream 20 Skim Milk 0	50.00 11.00	60.00 6.50	$70.00 \\ 0.50$				
5 Cream 50 Whole Milk 4	16.5 44.5	$20.50 \\ 46.00$	$\frac{24.50}{46.00}$	28.25 46.75	32.50 47.25		
6 Cream 40 Whole Milk 4	$21.0 \\ 40.0$	$26.00 \\ 40.50$	$\frac{31.25}{39.25}$	36.25 38.75	41 25 38.50		
7 Cream 30 Whole Milk 4	$\frac{29.0}{32.0}$	$\frac{36.00}{30.50}$	43.25 27.25	50.00 25.00	57.00 22.75		
8 Cream 20 Whole Milk 4	48.0 13.0	58 50 8.00	70.0 0.5				
Add to each above combination							
Unsweetened Condensed Skim Milk*	24.00	18.5 15.0	14.5 15.0	10.00 15.00	5.25 15.00		
Sugar	$15.0 \\ 0.3$	0.3	0.3	0.3	0.3		
Total	100.0	100.0	100.0	100.0	100.0		

^{*} Concentration, 3 to 1; contains 27% solids.

Note: The proportions given in columns a and b represent medium-fat ice creams commonly produced for soda fountain trade. These types of ice cream usually have a very smooth texture. The increased scrum solids are derived chiefly from concentrated milk products. In some cases about 90% of the serum solids are added in the form of condensed skim milk, which means that approximately one-third of the mixture is condensed milk and one-fifth is cream testing 40 per cent fat. The cream flavor may be largely masked by the condensed-milk flavor, particularly if the latter has a pronounced cooked flavor. Consequently, the flavor will be improved by using either whole or skim milk with a minimum quantity of condensed skim milk.

either whole or skim milk with a minimum quantity of condensed skim milk.

The proportions given in columns c, d, and c represent ice creams with smaller additions of serum solids in the form of condensed skim milk, than those shown in columns a and b. It is believed that a small addition of serum solids to the higher fat products will improve the original texture, and in preventing deterioration of

texture during storage.

Table 3.—Amounts of Cream of Different Fut Content, Dry Skim Milk, and Fresh Skim Milk or Whole Milk Necessary for Making Different Types of Ice Crean

Skiin Britte OF WHOIS MIII	k Necessary	for Makin	g Different	Types of	Ice Cream	
A. Solids	Types of Ice Cream					
	a	ь	o	d	a	
	Per Cent	Per Cent	Per Cent	Per Cent	Per Cent	
Fat	10	12	14	16	18	
Serum Solids	11	10	9	8	7	
Sugar	15	15	15	15	15	
Gelatin	0.3	0.3	0.3	0.3	0.3	
Total Solids	36.3	37.3	38.3	39.3	40.3	
B, Ingredients Per cent C.	. Percentage	of Ingred Ice (lient by W Cream Mixt	eight in E	ach Type of	
	a	ъ	c	d	e	
1 Cream 50	20.00	24.00	28.00	32.00	36.00	
Skim Milk 0	60.00	37.00	54.00	51.00	48.00	
2 Cream 40	25.00	30,00	35.00	40.00	45.00	
Skim Milk 0	55.00	51.00	47.00	43,00	39.00	
3 Cream 30	33.25	40.00	46.75	53.25	60.00	
Skim Milk 0	46.75	41.00	35.25	29.75	24.00	
4 Cream 20	50.00	60 00	70.00	80.00		
Skim Milk 0	30.00	21.00	12.00	3.00		
5 Cream 50	15.00	19.00	23.50	27.57	32.00	
Whole Milk 4	65.00	62.00	58.50	55.43	52.00	
6 Cream 40	19.00	24.50	30.00	35.25	40.75	
Whole Milk 4	61.00	56.50	52.00	47.75	43.25	
8 Cream 20	26.25	33.75	41.50	48.75	56.50	
Whole Milk 4	53.75	47.25	40.50	34 25	27.50	
7 Cream 30	42.50	54.75	67.00	79.25		
Whole Milk 4	37.50	26.25	15.00	3.75		
Add to each above com-						
bination	F 00	4.00	2.00	0.00	1.00	
Dry Skim Milk	5.00	4.00	3.00	2.00	1.00	
Sugar Gelatin	15.00 0.3	15 00 0.3	15 00 0.3	15.00 0.3	15.00 0.3	
m	100.00	100.00	100.00	100.00		
Total	100.00	100.00	100.00	T00*00	100.00	

Note: Dry skim milk is a very convenient form of serum solids to use in the manufacture of ice cream. Tests reported in U. S. Department of Agriculture Circular 179 have shown that the addition of dry skim milk will produce a medium grade ice cream equal to ice creams made with condensed milk. The principal criticisms of ice creams containing dry skim milk are usually due to the flavor imparted by this product. The formulas given in the above table will reduce this difficulty to a minimum by using as much whole and skim milk as possible in the preparation of the mixes.

No. 7			No. 8		
8% Fat-Cream, Skim		Milk	8% Fat-Sweet Butter,		
Powder, 14%	Sugar.		Skim Milk Powder, 14	% Bugar	۲.
Sugar	14	lb.	Sugar	14	lb.
Gelatin	0.5	lb.	Gelatin	0.5	lb.
Skim Milk Powder	6	lb.	Skim Milk Powder	6	lb.
Cream (25%)	20	lb.	Butter (84%)	6.1	lb.
Skim Milk	59.5	lb.	Mılk (4%)	73.4	lb.

Table 4.—Amounts of Cream of Different Fat Content, Skim Milk, and Sweetened Condensed Whole Milk Necessary for Making Different Types of Ice Cream

A. Solids	Types of Ice Cream				
	a	b	c	d	e
	Per Cent	Per Cent	Per Cent	Per Cent	Per Cent
Fat	10	12	14	16	18
Serum Solids	11	10	9	8	7
Sugar	15	15	15	15	15
Gelatin	0.3	0.3	0.3	0.3	0.3
Total Solids	36.3	37.3	38.3	39.3	40.3
B. Ingredients Per cent C.	Percentage of Ingredient by Weight in Each Type of Ice Cream Mixture				
	a	b	o	d	6
1 Cream 50	16,00	20.68	25.6	30.4	35.2
Skim Milk 0	50.50	49.72	49.2	48.8	46.9
2 Cream 40	20.00	26.00	32.0	38.0	44.25
Skim Milk 0	46.50	44.40	42.8	41.2	37.85
3 Cream 30	26.66	34.7	42.7	50.7	58.7
Skim Milk 0	39.84	35.70	32.1	28.5	23.4
4 Cream 20	40.00	52.0	64.0	76.0	
Skim Milk 0	26.50	18.4	10.8	3,2	
Add to each above com-					
bination					
Sweetened Condensed					
Whole Milk*	25.00	20.00	15.00	10.00	5.00
Sugar	4.5	6.6	8.7	10.8	12.9
Water	4.0	3.0	1.5	_	
Gelatin	0.3	0.3	0.3	0.3	0.3
Total	100.0	100.0	100.0	100.0	100.0
* Contains 8% fat, 23% s	erum solids	, and 42%	sugar.		

Note: Before using these formulas the manufacturer should be certain that the analysis of the sweetened condensed whole milk conforms to the analysis used in compiling this table,

lb. lb. lb.

8% Fat—Sweet Butter, Skim Milk, Skim Cream (25%) 28	lb.
Milk Powder, 14% Sugar. Milk (4%) 27.5	lb.
Sugar 14 lb. No. 12	
Gelatin 0.5 lb. 8% Fat—Cream, Milk, Sweet Conde	nsed
Daint Milk Lowder 0 10.	
1741101 (01/0)	11.
Guart Candonad Mills 00	lb.
by rat—sweet butter, water, Skill Comm (250)	
Milk Powder, 14% Sugar. Milk (4%) 53.7	
Sugar 14 lb. Milk (170)	
Gelatin 0.5 lb. No. 13	
Butter (84% 9.6 lb. 8% Fat—Sweet Butter, Skim Milk	650
Skim Milk Powder 12.0 lb. Sweet Condensed Skim Milk	and
Water 03.3 lb. 14% Sugar	
No. 11 Sugar 2.8	lb.
8% Fat-Cream and Milk, Condensed Gelatin 0.5	
	lb.
Sugar 14 lb. Sweet Butter 9.6	
Gelatin 0.5 lb. Skim Milk 59.1	lb.

Table 5.—Amounts of Cream with 50% of the Fat Added in the Form of Butter, Unsweetened Condensed Skim Milk, and Dry Skim Milk Necessary for Making Different Types of Ice Cream

A.	Solida		Types of Ice Cream				
			а	ь	c	d	c
			Per Cent	Per Cent	Per Cent	Per Cent	Per Cent
	t		10	12	14	16	18
	rum Solids		11	10	9	8	7
	gar		15	15	15	15	15
	latın		0.3	0.3	0.3	0.3	0.3
	Fotal Solids	• • • • •	36.3	37.3	38.3	39.3	40.3
В.	Ingredients P	ercent C	r cent C. Percentage of Ingredient by Weight in Each Type fat lee Cream Mixture				
			a	b	c	d	e
1	Cream Unsweetened	40	12.50	15.00	17.50	20.00	22.50
	Condensed		00.00	34 00	30.00	26 00	22.00
	Skim Milk* Water		28.00 28.40	28.68	28 96	29.25	29.53
_						20.00	22.50
2	Cream	40	12 50	15.00	17.50 8.51	7 60	6.25
	Dry Skim Milkt		10 80	9.67 53.01	50.45	47.65	45.28
	Water		55.60				
3	Cream Unsweetened	20	25.00	30.00	35.00	40.00	45.00
	Condensed Skim Milk*		34 00	29.00	24.00	19.00	14.00
	Water		19.00	18.68	17.46	16.25	15.03
		00		30.00	35.00	40.00	45.00
4	Cream	20	25.00	8.25	6.25	5.38	6.00
	Dry Skim Milkt		9.60 44.30	39.43	35.21	29.87	23.03
	Water		44.50	35.43		20.01	20.00
	d to each above	com-					
	ination	00	6.10	7.32	8 5 4	9.75	10.97
	tter	82	15.00	15 00	15.00	15.00	15.00
	gar		0.3	0.3	0.3	0.3	0.3
	latın				100.0	100.0	100.0
- 7	Гоtаl		100.0	100.0		100.0	100.0
	* Concentration	ratio, 3 f	to 1: contair	18 27% Bul	1d8.		

^{*} Concentration ratio, 3 to 1; contains 27% solids.

Note: In the preparation of ice cream mixes with butter only the freshest and best grades of unsalted butter should be used.

No. 14		No. 16	
8% Fat-Cream, Milk, Eva 14% Sugar.	porated Milk,	10% Fat—Cream, Skim M Powder, 12% S	ugar.
Sugar	14 lb.	Sugar	12 lb.
Gelatin	0.5 lb.	Gelatin	0.5 lb.
Evaporated Milk (8%)	30 lb.	Skim Milk Powder	5 lb.
Cream (25%)	16 lb.	Cream (25%)	40 lb.
Milk (4%)	39.5 lb.	Skim Milk	42.5 lb.
No. 15	00.0 101	No. 17	
10% Fat-Cream, Whole	Milk, Skim	10% Fat-Sweet Butter,	Whole Milk,
Milk Powder, 12%		Skim Milk Powder, 1:	
Sugar	12 lb.	Sugar	12 lb.
Gelatin	0.5 lb.	Gelatin	0.5 lb.
Skim Milk Powder	5 lb.	Skim Milk Powder	6 lb.
Cream (30%)	26 lb.	Butter (84%)	9 lb.
Milk (4%)	56.5 lb.	Milk (4%)	72.5 lb.

^{† 95%} solids.

Table 6.—Amounts of Cream of Different Fat Content, Unsweetened Condensed Skim Milk and Fresh Whole or Skim Milk Necessary for Making Different Types of Mixes Without Sugar

A. Solids		Туре	s of Ice C	ream	
	а	ь	o	d	e
	Per Cent	Per Cent	Per Cent	Per Cent	Per Cent
Fat	10	12	14	16	18
Serum Solids	11	10	9	8	7
Sugar	15	15	15	15	15
Gelatin	0.3	0.3	0.3	0.3	0.3
Total Solids	36.3	37.3	38.3	39.3	40.3
B. Ingredients Per cent C.	t C. Percentage of Ingredient by Weight in Each Ty				ach Type of
	а	b	o	d	e
1 Cream 50	23.50	28.25	33.00	37.75	42.50
Skim Milk 0	48.75	49.25	49.75	50.25	50.50
2 Cream 40	29.5	35,25	41.25	47.00	53.00
Skim Milk 0	42.75	42.25	41.50	41.00	40.00
3 Cream 30	39.25	47.00	55.00	62.75	70.5
Skim Milk 0	33,00	30.50	27.75	25.25	22.5
4 Cream 20	58.75	70.50	82.50		
Skim Milk 0	13.50	7.00	0.25		
5 Cream 50	19.25	24.00	28.75	33.25	38.00
Whole Milk 4	53. 00	53.50	54.00	54.75	55.00
6 Cream 40	24.75	30.75	36.75	42.50	48.50
Whole Milk 4	47.50	46.75	46.00	45.50	44.50
7 Cream 30	34.00	42.50	50.75	59.00	67.25
Whole Milk 4	38,25	35.00	32.00	29.00	25.75
8 Cream 20	55.50	69.00	82.25		
Whole Milk 4	16.75	8.50	0.50		
Add to each above com- bination					
Unsweetened Condensed Skim Milk*	27.75	22.50	17.25	12.00	
Gelatin	0.34	0.34	0.34	0.34	7.00 0.34
	100.0	100.0	100.0		–
Total	100.0	100.0	100.0	100.0	100.0

^{*}Concentration ratio, 3 to 1; contains 27% solids.

Note: Ice cream mixes made from the formulas in Tables 6 and 7 should not be confused with mixes containing sugar. For every 100 pounds of ice cream desired use 85 pounds of mix and add 15 pounds of sugar. In case the manufacturer desires to use 1 or 2 pounds more or less of sugar, the basic formulas will not be materially changed.

No. 18	1	No. 19	
10% Fat-Sweet Butter, Powder, Water, 12%		10% Fat—Sweet Butter, Powder, Water, 12%	Skim Milk Sugar.
Sugar	12 lb.	Sugar	12 lb.
Gelatin	0.5 lb.	Gelatin	0.5 lb.
Skim Milk Powder	6 lb.	Butter (84%)	12 lb.
Butter (84%)	12 lb.	Skim Milk Powder	12 lb.
Skim Milk	69.5 lb.	Water	63.5 lb.

Table 7.—Amounts of Cream of Different Fat Content, Dry Skim Milk and Fresh Skim or Whole Milk Necessary for Making Different Types of Mixes Without Sugar

	WILII	out sugar					
A. Solids		Type	es of Ice C				
	a	b	c c				
	Per Cent	Per Cent	Per Cent	d Dom Oomt	e DO4		
		1 er cent	rercent	Per Cent	Per Cent		
Fat	10	12	14	16	18		
Serum Solids		10	9	8	7		
Sugar		15	15	15	15		
Gelatin		0.3	0.3	0.3	0.3		
Total Bolids	36.3	37.3	38.3	39.3	40.3		
B. Ingredients Per cen	t C. Percentage	of Ingred	of Ingredient by Weight in Each Type of Ice Cream Mixture				
	a	b	c	d	r		
1 Cream 50	23.52	00.00	00.04				
Skim Milk 0	70.69	28.22	32.94	37.64	42.35		
2 Cream 40		66.14	63.47	60.04	56.38		
Skim Milk 0	29.40	35.27	41.17	47.05	52.92		
	64.81	60,09	55.24	50.63	45.81		
	39.20	47.03	54.90	62.73	70.57		
	55.01	48,33	41.51	34.95	28.16		
4 Cream 20	58.80	70.55	82.35	94.10			
Skim Milk 0	35.41	24.81	14.06	3.58			
5 Cream 50	17.40	22.36	27.50	32.50	37.50		
Whole Milk 4	76.81	73.00	68 91	65.18	61.23		
6 Cream 40	22.27	28.60	35 00	41.50	49.00		
Whole Milk 4	71.94	66.76	61.41	56.18	49.73		
7 Cream 30	30.77	39 60	48.50	57.50	66.25		
Whole Milk 4	63.44	55.76	47.91	40.18	32.48		
8 Cream 20	50.0	64.26	78 75	93.25			
Whole Milk 4	44.21	33.10	17.66	4.43	Westernament .		
Add to each above com-							
bination							
Dry Skim Milk*	5.79	4.64	3 59	2.32	1.27		
Gelatin	0.34	0.34	0.34	0.34	0.34		
Total	100.0	100 0	100.0	100.0	100.0		
*Contains 95% solids.							
No. 20		1	N	0. 22			
10% Fat-Cream, Whol	e Milk Skim	100% F	at—Sweet		hala Min		
Milk Powder, 14%	Sugar.	Skii	m Milk Po	wder, 14%	Sugar.		
Sugar Gelatin	14 lb. 0.5 lb.	Sugar Gelati			14 lb.		
Skim Milk Powder	0.5 lb. 4 lb.		n Milk Powd		0.5 1ь.		
Cream (30%)	26 lb.		мик гоwа г (84%)	+·F	4 lb. 9 lb.		
Milk (4%)	55.5 lb.	Milk			9 lb. 72.5 lb.		
No. 21	00.0			o. 23	12.0 10.		
10% Fat—Cream, Skim M Powder, 14% S	ilk, Skim Milk	10% I Skir	at—Sweet n Milk Por	Butter, Sl	cim Milk,		
Sugar	14 lb.	Sugar		, 11/0	14 lb.		
Gelatin	0.5 lb.	Gelatn	n		0.5 lb.		
Skim Milk Powder	4 lb.		 Milk Powd	er	4 lb.		
Cream (25%)	40 lb.		Butter (8		12 lb.		
Skim Milk	41.5 lb.	Skim	Milk		69.5 lb.		

		I		
No. 24		No. 31		
10% Fat-Sweet Butter, Skim Milk Pow- der, Water, 14% Sugar.		12% Fat—Sweet Butter, Whole Milk, Skim Milk Powder, 12% Sugar.		
Sugar	14 lb.	Sugar	12 lb.	
Gelatin	0.5 lb.	Gelatin	0.5 lb.	
Butter (84%)	12 lb.	Skun Milk Powder	3 lb.	
Skim Milk Powder	10.6 lb.	Butter (84%)	10.8 lb.	
Water	62.9 lb.	Milk (4%)	73.7 lb.	
No. 25	02.0	No. 32	10.1 10.	
10% Fat-Cream, Milk Cor		12% Fat-Sweet Butter, Skim Milk and		
Milk (27%), 14% S		Skim Milk Powder, 12		
Sugar	14 lb.	Sugar	12 lb.	
Gelatin	0.5 lb.	Gelatin	0.5 lb.	
Condensed Milk	18 lb. 28 lb.	Skim Milk Powder	3 lb.	
Cream (30%) Milk (4%)	28 lb. 39.5 lb.	Butter (84%)	14.3 lb.	
	39.3 10.	Skim Milk	70.2 lb.	
No. 26		No. 33		
10% Fat—Cream and Milk, Sweet Con- densed Whole Milk, 14% Sugar.		12% Fat—Sweet Butter, Sk der, Water, 12% S	im Milk Pow- ugar.	
Sugar	6.8 lb.	Sugar	12 lb.	
Gelatin	05 lb.	Gelatin	0.5 lb.	
Condensed Milk	18 lb.	Butter (84%) Skim Milk Powder	14.3 lb.	
Cream (25%)	27 lb.	Skim Milk Powder	9.5 lb.	
Milk (4%)	47.7 lb.	Water	63.7 lb.	
No. 27		No. 34		
10% Fat—Sweet Butter, Sk	im Milk and	12% Fat-Cream, Whole	Milk. Skim	
Sweet Condensed Skin	n Milk,	Milk Powder, 14%	Sugar.	
14% Sugar.		Sugar	14 lb.	
Sugar	14 lb.	Gelatin	0.5 lb.	
Gelatin	0.5 lb.	Skim Milk Powder	2 lb.	
Butter (84%)	12 lb.	Cream (25%)	41.5 lb.	
Condensed Skim Milk	16 lb.	Milk (4%)	42 lb.	
Skim Milk	57.5 lb.	No. 35		
No. 28				
10% Fat-Cream, Milk Eva	porated Milk,	12% Fat—Cream, Skim M 14% Sugar.	,	
Sugar	14 lb.	Sugar	14 lb.	
Gelatin	0.5 lb.	Gelatin	0.5 lb.	
Evaporated Milk	18 lb.	Skun Milk Powder	2 lb.	
Cream (25%)	28 lb.	Cream (30%)	40 lb.	
Milk (4%)	39.5 1Ь.	Skim Milk	43.5 lb.	
No. 29		No. 36		
12% Fat—Cream, Whole Milk Powder, 12% S	Milk, Skim ugar.	12% Fat—Sweet Butter, Skim Milk Powder, 14	Whole Milk, % Sugar.	
Sugar	12 lb.	Sugar	14 lb.	
Gelatin	0.5 lb.	Gelatin	0.5 lb.	
Skim Milk Powder	2 lb.	Skim Milk Powder	2.5 lb.	
Cream (25%)	41 lb.	Butter (84%)	11 lb.	
Milk (4%)	44 lb.	Milk (4%)	72 lb.	
No. 30		No. 37		
12% Fat—Cream, Skim I Milk Powder, 12% S		12% Fat—Sweet Butter, Skim Milk Powder, 149	Skim Milk, % Sugar.	
Sugar	12 lb.	Sugar	14 lb.	
Gelatin	0.5 lb.	Gelatin	0.5 lb.	
Skim Milk	3 lb.	Skim Milk Powder	2.5 lb.	
Cream (30%)	40 lb.	Butter (84%)	14.3 lb.	
Skim Milk	44.5 lb.	Skim Milk	68.7 lb.	

FOOD P	RODUCTS, B	EVERAGES, FLAVORS	143
No. 38		No. 45	
12% Fat-Sweet Butter, Skim Milk Pow- der, Water, 14% Sugar.		14% Fut-Sweet Butter Skim Milk Powder, 1	r, Whole Milk,
Sugar	14 lb.	Sugar	
Gelatin	0.5 lb.	Gelatin	12 lb, 0.5 lb,
Butter (84%)	14 3 lb.	Skim Milk Powder	
Skim Milk Powder	9 lb.		
Water	62.2 lb.	Butter (84%) Milk (4%)	13.3 lb 72.2 lb.
No. 39	02.2 ID	No. 46	12.2 10.
12% Fat—Cream, Milk, Co Milk (27%), 11% S		1407 Fnt-Sweet Butter	
Sugar	14 lb.	Skim Milk Powder, 1	
Gelatin	0.5 lb.	Sugar	12 lb.
		Gelatin	0.5 lb.
Condensed Milk	16 lb.	Skim Milk Powder	2 lb.
Cream (30%)	35.5 lb.	Butter (84%)	16 7 lb.
Milk (4%)	34 lb.	Skun Milk	68.8 lb.
No. 40	0 . 1 1	No. 47	
12% Fat—Cream, Milk. Swe Whole Milk, 14%	Sugar.	14% Fat—Sweet Butte Powder, Water, 129	
Sugar	8 1 lb.	Sugar	12 lb.
Gelatin	0.5 lb.	Gelatm	0.5 lb.
Sweet Condensed Milk	14 lb,	Butter (84%)	16.7 lb.
Cream (25%)	38 lb.	Skim Milk Powder	
Milk (4%)	39.1 lb.	Water	8.6 lb.
No. 41			62.2 lb.
12% Fat—Sweet Butter, Skim Milk and Sweet Condensed Skim Milk, 14% Sugar.		No. 48 147 FatCream, Milk, Skim Milk Powder, 14% Sugar.	
_		Sugar	.,
Bugar	9.2 lb.		
Gelatin	0.5 lb.	Gelatin	0.5 16.
Sweet Skim Condensed M		Skin Milk Powder	1 lb.
Butter (84%)	14.3 lb.	Cream (30%)	41 lb.
Skim Milk	64 lb.	Milk (4%)	43.5 lb.
No. 42		No. 49	
12% Fat—Cream, Milk, 1 Milk, 14% Suga		14% Fat- Cream, Skim Milk Powder, 14%	Milk, Skim Sugar.
Sugar	14 lb	Sugar	14 lb.
Gelatin	0.5 lb.	Gelatin	0.5 lb.
Evaporated Milk (8%)	20 lb.	Skim Milk Powder	1 lb.
Cream (30%)	30 lb.	Cream (25%)	56 lb.
Milk (4%)	35.5 lb.	Skim Milk	28.5 1ь.
No. 43		No. 50	
14% Fat-Cream, Whole Milk Powder, 12% S		14% Fat—Sweet Butter, Skim Milk Powder, 1-	
Sugar	12 lb.	Sugar	14 lb.
Gelatin	0.5 lb.	Gelictin	0.5 lb.
Skim Milk Powder	2 lb.	Skim Milk Powder	1.1 lb.
Cream (30%)	14 lb.	Butter (84%)	13.3 lb.
Milk (4%)	44.5 lb.	Milk (4%)	71.1 lb.
No. 44	l	No. 51	
4% Fat-Cream, Skim Mill Powder, 12% Sug		14% Fat—Sweet Butter, Skim Milk Powder, 1s	
	12 lb.		14 lb.
Sugar	0.5 lb.	Bugar Gelatin	
Sugar Gelatin	0.5 lb.	Gelatin	0.5 lb.
Sugar			

No. 52	
14% Fat-Sweet Butter, 8	kim Milk Pow
der, Water, 14%	Sugar.
Sugar	14 lb.
Gelatin	0.5 lb.
Butter (84%)	16.7 lb.
Skim Milk Powder	7.6 lb.
Water	61.2 lb.
No. 53	
14% Fat-Cream, Milk, Co	ondensed Skim
Milk (27%), 14%	
Sugar	14 lb.
Gelatin	0.5 lb.
Condensed Skim Milk	6 lb.
Cream (30%)	42 lb.
Milk (4%)	37.5 lb.
No. 54	
14% Fat-Cream, Milk, Sw	root Condonnad
Whole Milk, 14%	Sugar.
Sugar	11.6 lb.
(lelatin	0.5 lb.
Sweet Condensed Milk (8	3%) 6 lb.
Cream (38%)	40 lb.
Milk (4%)	41.9 lb.
No. 55	
14% Fat-Sweet Butter, S	kim Milk and
Sweet Condensed Ski	m Milk.
14% Sugar.	
Sumr	11 6 lb

11/0 Dugai.	
Sugar	11.6 lb.
Gelatin	0.5 lb.
Condensed Skim Milk	6 lb.
Butter (84%)	16.7 lb.
Skim Milk	65.2 lb.
No. 56	
14% Fat-Cream, Milk,	Evaporated

14% Fat—Cream, Milk, Evaporated Milk, 14% Sugar.

Bugar	14 lb.
Gelatin	0.5 lb.
Evaporated Milk	10 lb.
Cream (30%)	40 lb.
Milk (4%)	25.5 lb

Fig Cream

For a 10-gal, finished ice cream.

45 lb. unflavored mix No. 10 can of solid packed pie figs ground fine in a food chopper is added while the mix is in the freezer.

Fig and Walnut Ice Cream

For a 10-gal. batch of finished product take 3 lb. canned pie figs, 2 lb. walnuts, run them through the fruit chopper, not too fine, and add the same as for strawberries. Use either English or black walnuts. The English are rather high in price.

The gelatin given in these formulas are .5 of a pound of high grade gelatin,

or you may use half good ice cream powder and half gelatin.

When mix is ready pasteurize the whole mix at 145 to 150° F., then viscolize or homogenize the whole mix while hot; cool to 40 or 50° F., age for 24 to 48 hours, then freeze.

Simple Ice Cream Mix

Cream (30%)	35.8 lb.
Milk (3.5%)	49.7 lb.
Sugar	14 lb.
Gelatin	0.5 lb.
100.0 lb. of mix containing and 33.4% total solids.	12.5% fat

Complex Ice Cream Mix

Cream (30%)	41.7 lb.
Condensed Skim Milk	15.3 lb.
Skim Mılk	28.5 lb.
Sugar	14 lb.
Gelatin	0.5 lb.
100.0 lb. of mix containing and 37% total solids. A standard vanilla extract to	dd 91/4 oz
of mix.	

Preparing 20% Cream

To make 360 lb. of 20% cream use 160 lb. of 40% cream and 200 lb. of 4% milk.

Preparing 35% Cream

To make 360 lb. of 35% cream use 310 lb. of 40% cream and 50 lb. of 4% milk.

Chocolate Ice Cream

	CHUCK	natu	100	Crean	OL.	
Milk					32	oz.
Sugar					16	OZ.
Flour					2	OZ.
Salt					-16	oz.
Eggs					4′°	oz.
Cream					32	oz.
Vanillin	1				1/4	OZ.
Unswee	tened	Cho	colat	A	4~	07

Heat milk and add flour, salt, and sugar. Stir thoroughly in double boiler for 20 minutes after batch is brought to a boil. After the mass thickens, add the beaten eggs and cook for 5 minutes longer with constant stirring. Cool, add cream which has been whipped into a stiff paste, and then add the flavoring. Add the melted chocolate, previously mixed with a litle sugar and warm milk to form a paste. Put in a refrigerator or pack in ice and salt until frozen.

Ice Cream Without Gelat

Butter Fat	12	lb.
Sugar (Granulated)	12	lb.
Cerelose (Corn Sugar)	4	lb.
Milk Serum Solids Vanilla Flavor		5 lb. suit

Preventing Sandiness in Ice Cream U. S. Patent 1.940.109

By freezing and whipping air into icecream nux at such a rate that 30% of the water is frozen in less than 1 minute a smoother product than usual is obtained and one in which the milk solids may be increased with less likelihood of forming "sandy" ice cream.

Water Ices and Sherbets

The figures are given on the basis of 100 lb, of mix which is about 101/2 gal.

Water Ico

Cane Sugar	25 10.
Corn Sugar	7 lb.
Agar (3.2 oz. or 90.6 g.)	0.2 lb.
Gum Tragacanth or Ga-	
lagum C (6.4 oz. or	
181.2 g.)	0.4 lb.
Water, Fruit, Fruit Acid,	

67.4 lb. Flavor, and Color

Overrun 20 to 25%. Total yield 13 gal.

Sherbet Using Milk

Cane Sugar	25 lb.
Corn Sugar	7 lb.
Agar (3.2 oz. or 96.6 g.)	0.2 lb.
Gum Tragacanth or Ga-	
lagum C (3.2 oz. or	
90.6 g.)	0 2 lb.
Whole Milk	50 lb.
Title A There's There's Aniel	

Water, Fruit, Fruit Acid, Flavor, and Color 17.6 lb. Overrun 25 to 30%. Total yield 13.5 gal.

Sherbet Using Cream Mix

Cane Sugar

lb.

Agar (3.2 oz. or 90.6 g.) Gum Tragacanth or Ga-	0 2 lb.
lagum C (3.2 oz. or 90.6 g.)	0.2 lb.
Ice Cream Mix, without Sugar or Gelatin	10 lb.
Water, Fruit, Fruit Acid, Flavor, and Color	57.6 lb.

Overrun 25 to 30%. Total yield 13.5 gal.

Flavor, and Color

Orange Water Ice (For 10 Gal. Batch)

Granulated Sugar	21	lb.
Corn Sugar	7	lb.
Galagum C	3	OZ.
Orange Juice (or	Its Equiv-	
alent in Orange	Flavor) 1	gal.
Citric acid to suit.		10 gal
th water Takes p	o overrun.	

Orange Sherbet (10 Gal. Mix)

(10 (10)		
Cane Sugar	2214	lb.
Cerelose (Corn Sugar)	71/2	
Milk	4	gal.
Gelatin	11	07.,
Orange Concentrate	4	OZ.
	:4	Mak

Citric acid and color to suit. Make up to 10 gal, with water.

Corne Tunket

COCOR DUILAC		
Cocoa	2	oz.
Boiling Water	4	oz.
Sugar	4	OZ.
Milk	32	uz.
Junket Tablets	2	
Cold Water	1	OZ.
Vanilla Extract	1/4	OZ.

Cook mixture of cocoa and water in double boiler for five minutes. Add sugar, stir until dissolved, and then add milk which has been previously pre-heated to 100° F. Add vanilla extract and heat to 120° F. Stir in junket tablets which are dissolved first in a little water. Pour into containers immediately, let stand until set.

Reworking Cream

For cream of poor quality mix equal parts of the cream and water and heat to 135° F. in a fore warmer. Condense in a vacuum pan until a volume equal to that of the original cream is obtained. 1 se 3 parts of cream to 1 part of water for cream that is of a slightly higher grade but that has off flavors and odors. In this case fore warm and condense also until a volume equal to that of the original cream is obtained.

Composition of Mixes to Be Used in the Manufacture of Sweet Cream Cream Cheese

The most desirable cream cheese that has been manufactured by this method contains from 15 to 18% of dry skim milk and 20% of butterfat in the final cheese mix.

The following mixes will make a very desirable cream cheese:

Formula No. 1

Per handred pounds: lb. 20 Butterfat 15 lb. Dry Skim Milk 0.4 lb. (Ichtin (250 Bloom Test) 0.75 lb. Salt

Starter, 3 lb. (if cheese is for immediate consumption or 1 lb, if it is to be held in storage from 7 to 10 days prior to delivery to the consumer).

No. 2

Per hundred pounds: 20 lb. Butterfat Dry Skim Milk 18 lb. Gelatin (250 Bloom Test) 0 t lb. Sult 0.75 lb. Starter 3 or 1 lb. (as stated in No. 1)

No. 3

Per hundred pounds: lb. Batterfat Dry Skim Milk 15 lb. Sult 0.75 lb. Gelatin (250 Bloom Test) 0.4 lb. Starter, 3 or 1 lb, as stated in No. 1. It requires 7 to 10 days for a desirable

mild acid flavor to develop in the cream cheese when only 1 lb. of starter is used in the cheese mixes. However, 3 lb. of starter is sufficient to develop the desired acidity by the end of the second day, providing a high quality starter is used in the cheese. If the cheese is to be held in storage for a period of approxi-mately 30 days, 1 lb. of starter or a fraction thereof will develop the desired

oughly sterilized prior to use and all ingredients must be of high quality. The most desirable cream cheese is obtained when using No. 2, however either No. 1 or No. 3 furnishes a very

desirable cream cheese.

flavor. All equipment should be thor-

The addition of dry skim milk, starter, salt and gelatin reduces the butterfat content of the resultant mix and sufficient fat must be added to the mix to replace the decrease in butterfat content by the addition of these ingredients. The addition of 1% of dry skim milk and other non-fat ingredients reduces the butterfat content of the finished cheese mix 0.26 of 1%. Therefore, in preparing a mixture that will furnish a butterfat content of 20% in the finished cheese when using 15% of dry skim milk, the cream from which the cheese is to be made must test 23.9% butterfat.

Cream Cheese (Geneva Method)

(Detailed Directions for 100-lb. Batches) Acid Flavor

Add 5 lb. of dry skim milk to 93 lb. of sweet cream testing 40 to 42% milk fat. Then add 0.5 lb. of ground agar and 0.75 lb. of salt. The cream should be well agitated as the dry skim milk and agar are slowly added. Pasteurize at 180° to 185° F. for 5 minutes. Cool to 110° F. Add 0.75 lb. of commercial starter. Homogenize at 3500 lb. pressure using no strainer in the intake pipe line. The homogenizer should have been previously run with water at 160° F. or above. Place the cheese immediately into the final package. Chill in a refrigerator at 10° to a temperature of 70° F. and incubate for 12 to 24 hours to develop an acid flavor. Then chill to and hold at 40° F.

The acidity develops slowly and the rate of development is controlled by the percentage inoculation. Reducing the skan milk solids to 3% tends to soften the body of the cheese and mereases the tendency towards some whey dramage and lower total acidity. The cheese may be softened by decreasing the homogenization pressure to 3000 lb, or firmed by increasing it to 4000 lb. More than 1 lb. of salt will retard and 11/2 lb. almost check acid development. Cream color umy be added before pasteurization, if desired, and it has the special advantage of reducing the intensification of color of cheese exposed to the air.

Consideration has also been given to the omission of starter and the securing of the desired acid flavor from Neuf-chatel, cottage, or Neufchatel cream The process itself presented no cheese. special difficulties (even cottage cheese could be homogenized in the cold or warm cream at 100 lb. pressure) and the mix-ture was treated in the regular way. About 50% of these acid cheeses is required to impart a very mild acid flavor to the finished product; or a product such as that made from an enriched milk by the cottage cheese process could be homogenized alone. The process is somewhat complicated and the flavor of the finished cheese is very mild, but it has excellent keeping quality.

The homogenizer may be a source of microbial contamination and may chill the first material passing through it. For these reasons the hot water rinse just before use is always essential. The cream mixture was strained through a coarse strainer with approximately 1/16 inch openings and the strainer to the homogenizer was always removed from the pape line to permit an even flow of the cream mixture. Short pipe lines are very desirable to reduce mechanical losses.

The hot cheese may be transferred with a filling machine or by land to 3-or 5-lb. lined boxes for bulk sale. The usual mayonnaise jar filling machine can be used for filling jars, but some difficulty may be encountered in making the small tin foil or cellophane wrapped 1 to 4-oz, packages. These puckages are made from the cold cheese by mobiling into proper size with a machine or by cutting into the proper size with a machine or by cutting into the proper size with a machine or by cutting into the proper size with a remodeled butter cutter. Some ingenuity must be used in the details of placing the cheese in the package.

Ripened Cheese Flavor (Cheddar and Roquefort)

Add 5 lb, of dry skim milk to 69 25 lb, of sweet cream testing 40 to 12°, or fat. Then add 0.75 lb, of common salt. (The agair is not essential in this cheese, but it improves sheing qualities.) The cream should be well agitated as the dry skim milk is slowly added. Remove paratha, inchesceloth, or other coating from the surface of 25 lb, of well ripened American cheddar cheese and grind or slee the cheese. Cheese color appears to be de simble for cream cheese of the cheddar through the cream the usual cheddar cheese color.

For Roquefort flavor use 79.25 lb, of sweet cream, 5 lb, of dry skim milk, 15 lb, of Roquefort cheese and 0.75 lb, of common salt. The entire mixture should be pasteurized at 160° or at 180° F for numbers, depending upon the keeping quality desired. Homogenize at 2500 lb, pressure, the muchine having been previously run with hot water. Place the hot cheese directly into the final package and immediately store at 35° to 40° F in the refrigerator.

Less Roquefort cheese is generally required as a flavor than is the case for American cheddar. Many persons who object to the flavor of Roquefort cheese consume large helpings of Roquefort cheese may be used, but investigations have been limited to the two varieties of cheese mean the constraints of the constraints

The ripened cheeses readily soften and disperse in the cream when the temperature exceeds 145° F. No necessity of using an emilsifying saft was ever cu-countered, but tests demonstrated that these salts, such as disodium phosphate and sodium citrate, could be used in

limited amounts without interfering with the process.

Other Food Flavors

Coarsely ground sweet pickle relish (outon thivor is undesirable), puntento, ohve and nut, pincapple, and other food flavors may be used. Add 5 lb, of dry skim milk, 0.5 lb, of ground agar, and 0.75 lb, of salt to 73.5 lb, of creim testing 40 to 42% of fat. The cream should be well agitated as the dry skim milk and agar are added. Pastenrize at 180° to 185° P, for 5 munites. Homogenize at 3500 lb, pressaue, the machine having been pieviously rim with hot water. Stir the flavoring material, 20 lb, is about right for most foods, directly into the hot cheese. Place in the final package and store immediately in the refrigerator at 55° to 40° E.

In some instances there may be an excessive quantity of junce. This can be mixed in the cream just before homogenization, but if the acidity of the junce is high the cream mixture may be previously cooled to 120° to 140° P. before adding the junce and the homogenization pressure techned to prevent excessive fat changing and coagulation. If the hody is somewhat soft the dry skini milk may be mercessed to 74b.

Most finit flavors did not blend well with cream choose, but fart spicy flavors are generally satisfactory.

O, and N. Cream Cheese (Marquardt)

Standard ze ridk to 10% of fat, then pesternize at 160° F. for 30 minutes; and homogenize at 2500 lb. pressure and at 120° F.

Cool the batch to 72° F., and add 6.2% of commercial starter and 15 cc. of reanity to 1000 lb. of milk. On the following day dram and salt as in the ninking of old style cream cheese and analyze for fat.

May the cheese prepared in the above manner with 40% cream to obtain the desired cheese fat content. This may be 27, 30, 35 or 40%. Then add 0.1% of gum and 5% of 40% sour cream. Add enough saft to have 0.75% in the finished cheese. Heat this entire mixture to 140 of 11 and homogenize at 120° F, and 3000 lb. pressure.

Bel Pacse Cheese (Farrar)

Use raw milk containing 3 to 4% of fat. Add ½% of lactic culture, and an equal amount of 8, thermophilus culture when available. Set the milk at 107° F, with rennet at the rate of 8 oz. per 1000

lb. of milk. The curd is cut after 15 minutes. Then part of the whey is drawn, and the cheese curd is dipped rapidly into the molds.

The cheese should drain on reed mats for 6 hours, being turned frequently. It is desirable to have the room at 80° F. The cheese can be unde in brick molds or circular ones 8 inches in diameter. The cheese should be of a thickness when finished so that it will weigh 3 to 5 lb.

The cheeses are salted by submerging in 20% salt brine at 50 to 60° F. for 18 to 24 hours.

The cheeses after drying are placed in a curing room at 40° F, with a relative humidity ranging from 85 to 90° F.

After curing the cheeses are wrapped and pucked so as to avoid evaporation. This is exceedingly important. The cheeses cure in 6 to 12 weeks, depending upon the quality of the milk used.

Semi-Soft Cheese (Marquardt)

Use raw or pasteurized milk testing 3.5% in fat. Use I oz. cheese color per 1000 lb. of milk. Then udd 14% of commercial lactic enlture and 14% of S. helveticus culture and heat to 87° F. In about 2 hours the acid will increase .02 to .04 in the milk. Then dulate 8 oz. of rennet in cold water and add at this rate for each 1000 lb. of milk.

The milk should set for 30 minutes, and, 30 minutes after cutting it is dipped rapidly into brick or round molds. It is pressed with 10 lb. pressure for 8 hours.

After 24 hours the cheese is rubbed lightly with salt, and then placed in a brine for 24 to 48 hours. The brine is made by dissolving 18 lb. of salt in 82 lb. of water.

The cheese is cured at 53-57° F. for a short time, about 3 weeks. It is then placed in storage at 40° F.

Each cheese should weigh from 3 to 7 lb.

Walter Price Rapid Cottage Cheese Method

Pasteurize skim milk. Cool to 90° F. and add 5% of culture. Acid development of 0.5% will require only 5 hours. Finish making cheese according to standard procedure.

Note: Setting at 72 to 85° F. requires 12 to 18 hours for 0.5% of acid to develop.

Propagating lactic culture:

Select good grade of skim milk. Pasteurize to 180° F. for 1 honr. Cool to 72° F. Add 1% of culture from another

eulture. Incubate at 72° F. for 12 hours. Place in 40° F. room until ready for use.

Selecting natural culture:

Place 6 qts. of raw skim milk into a 72° F, inculator. After 12 hours select those having a firm curd. Select of the firm curd samples the one having best flavor. Use this as a propagating culture for future batches. Always inoculate from a day old culture.

Developing a commercial culture (Strep. Lact.):

Pasteurize skim milk to 180° F. for 1 hour in quart bottles. Cool to 72° F. and add a few drops of culture from a commercial culture. Incubate for 12 hours. Repeat pusteurization of a fresh batch of skim milk; and inoculate 1% from above culture. Repeat for 3 days, always using the culture just previously developed. After this period the culture is ready for use in cheese, butter, or cultured milk manufacture. Cultures should be transferred duily, and used for 3 weeks or a shorter period.

Developing Special Cultures (Bac. Bulgarieus of Lacto bacillus Acidophilus).

Follow above procedure for commercial cultures.

Incubate at 98° F.

Goats' Milk Cheese

Heat fresh milk to 88° F. Add 25 defore adding rennet dilute it in 20 times its volume of water. Cut in cubes 1 m. square after 45 mmutes. Allow to stand for 5 minutes, then dip into molds after stirring gently for 5 additional minutes.

The forms are made of 3X tin; they are 4½ in. in diameter, and 5 in. high. Each form has 5 rows of holes, the holes being 1 in. apart and ½ in. in diameter.

The cheese curd is not disturbed until it is sufficiently matted. It is then turned frequently. It remains in the hoops for 30 hours at 70° F. It is then rubbed with salt and placed in a curing room at 60° F. with a high humidity. The cheese should be wiped freely and turned. After 6 weeks they are ready to package. Each cheese wighs ½ lb, and requires 4½ lb, of milk. The cheese is white and has an agreeable flavor at 6 to 10 weeks.

Hokah Sage Cheese

To 6914 lb. of 40% fat content cream add 5 lb. of dry skim milk. Then add 1/4 lb. of common salt and a like amount

of agar agar (ground or powdered). Slice and grind 25 lb. of well cured cheddar cheese into the mixture and stir while heating the batch to 160 to 180° F. Hold at this temperature for 2 minutes and cool to 140° F. Then add 1 to 3 ce. of oil of Sage, Dalmatian. It should be diluted in a pint of water and then mixed into a gallon of the cheese mixture which in turn is mixed into the entire batch. The mixture is then homogenized at 3500 lb. pressure, the machine having been previously run with hot water. If the minimum amount of sage oil is used 1/2 oz. of sage leaves, Salva officinalis, may be added to the batch after homogenization. In using the leaves great care must be exercised in pulverizing them and removing stems and coarse leaves. Thorough incorporation is an essential. Extensive trials have indicated the desirability of using the oil of sage only.

The cheese should be packaged while hot, and stored at 35 to 40° F.

| Cheese Pikante (Marquardt Method) | Roquefort Cheese | 20 | lb. | Cheddar Cheese | 20 | lb. | Camembert Cheese | 20 | lb. | Salt | ¼ | lb. |

Add small quantities of black pepper, cayenne pepper, paprika, and grind through a fine grinder. The addition of 2 to 4% of Sauterne Wine improves the Pikante. Grind with products at 70° F, package and store at 52 to 40° F.

New York Style Sage Cheese

The regular method for making cheddar cheese is followed. At the start 100 bb, of milk for colored curd is used for each 1000 th, of milk. The small butch of milk is colored green. Both batches are made alike. At cheddaring time the curds of both batches are mixed and matted. Before pressing oil of sage, balmatian is atomized over the curd at the rate of ½ oz. per 1000 lb. of milk used.

The green color is prepared by soaking green corn, green oats, or alfalfa in water, grinding, and pressing in a cider press. The color must be prepared fresh each day. The amount to add to the small batch of milk depends upon the intensity of color desired.

Some manufacturers prefer to add the oil of sage to the milk before making the cheese.

The above method appears to be the one most commonly used. Other methods

have been described but produce less satisfactory results.

Ricotta Cheese (Marquardt)

Heat whey to 190° F, as it is drawn from the cheese vat. Then add sour whey until albumn flakes are like snowflakes. Stop heating when albumn collects on top of whey. Drain in molds or bugs. The cheese after draining is surface salted and ready for use.

The sour whey used should have 1% of acid. It may require a Bulgaricus culture to ncheve this. To flake out the albumin about 10% of sour whey must be added to the sweet whey. When whey only is used and drained in bugs the cheese is called negette.

Commonly 10% of skim milk is added to the sweet whey to increase the yields. Hoops used as molds should be 5 inches in dumeter and 9 inches high and perforated. If the molds are completely filled with moist cheese with a strainer dipper the cheese resulting will be 7 mehes high. The cheese is rubbed with salt and returned to the hoops for 2 hours after the draining period over night without pressure. The cheese should be dried in a room at 110° F, and wrapped in paper and placed in storage.

Maroni Cheese (Marquardt)

This is made by using the Ricotta method substituting whole milk for skim milk and adding 1977. It is molded in hoops 8 mehes in diameter and 10 mehes high, giving a flushed cheese 7 inches high. Ricotta Gras is also the name for the whole milk-whey combination.

Sapsago Cheese

This cheese is made principally in Glarus, Switzerland, from sour, skim milk of cows. It is known also as Schabzieger, Glarnerkase, and Krauterkase. It is chained to have been made in the thirteenth century; the authentic history at least dates back to the fifteenth century. Sapsage is a small, hard, green cheese flavored with the leaves of a species of clover; it is shaped like a truncated cone, 4 inches high, 3 inches at the top. This cheese is imported to some extent into the United States under the name of Sap Sago.

The skim milk from which this cheese is made is not allowed to become sour enough to coagulate on heating, as it would make too hard a curd. The milk, when it has reached the right acidity, is heated to the boiling temperature while being stirred. Cold buttermilk is then added, as is also some whey having a high percentage of acidity. The material coagulating on the surface is skimmed off. The milk is then stirred, while sufficient acid whey is added to precipitate the casein. When too little whey is used the curd is too soft, and when too much is used it is too hard. The curd is dipped with a skimmer and spread out to cool and then put into boxes and allowed to drain and ferment. The box is kept at a temperature of above 60° F., and pressure is applied by weighting with stones. Ripening is allowed to continue from three to six weeks. If the temperature of the room is too high or if sufficient pressure is not applied, too rapid and strong fermentation results. The curd is used for making the finished product, but the cheese is seldom finished where the curd is made. The curd is ground in a mill, and for every 100 lb. of cheese there is added 5 lb. of salt and 25 lb. of dried Mehlotus caernlea, un aromatic clover which is grown in the Canton of Schweiz for the purpose. The ground material is worked up into a dough and is forced into molds lined with linen cloth and the name of the manufacturer is stamped on the large end. The mold is then emptied and re-filled. The cheeses are dumped promisenonsly into a large cask holding about 200 lb. A comparatively small quantity is shipped rato this country. It sells at a low price and is usually grated.

Red Cheese Rind Color

Formula No. 1

Sudan 4 dye is dissolved in equal parts of 70% alcohol and acetone, or

No.

Tonrnesol, Fachsin, or Bordeux Red dissolved in water (distilled water is preferred), or

No. 3

Iron Oxide, known also as Berlin Red or English Red made into a paste with a heavy oil.

The intensity of the color can be varied by changing the amount of the coloring substance.

Apply to outside of cheese.

Cheese, Ice-Cream and Salad Stabilizer U. S. Patent 2.007.218

0. 0. 1 110010 2,001,210		
Locust Bean Gum	65	oz.
Irish Moss, Powdered	35	oz.
Karaya Gum	15	oz.

When used in the preparation of cream cheese, the undiluted mixture of the three ingredients mentioned above is added at the time that the curds are mixed with the cream in the usual procedure for the manufacture of cream cheese, and in the proportion of about one-half of 1% by weight on a wet basis. The material is hented to about 165° F., homogenized, and then packed hot.

In ice cream it is used diluted with sugar, in the preferred proportion of one-half of 1% on a wet basis, the stabilizer acts to prevent crystallization of ice particles and thus insures a fine, smooth texture and a body which will hold up under severe shocks, such as are encountered in transportation and handhing. The use of it in ree cream also usually results in more rapid freezing, especially in old-style freezers.

Cheese Emulsifiers

U. S. Patent 1,940,031

1-4% of either of following are used: Sodium Mucate Sodium Lactate

Preservation of Rindless Cheese British Patent 434,374

Bacterial action on surface of rindless cheese is prevented by treatment with following prior to heating to 65° C.

Hydrogen Peroxide (35%) 0.3%

Low fat content cheese is heated to 65° C. The peroxide is added, mixed and later heated to 80° C.

Brandy Cheese

Use regular cheddar cheese, preferably an entire small cheese with the surfaces scruped clean, and allow to dry at room temperature for 2 to 4 weeks. Then place cheese in clear water at 40 to 80° F, for several days.

The cheeses are then placed in a mixture of brandy and high grade vinegar for several days. The brandy may be mixed in equal parts or less with the vinegar. Three per cent of salt should be added with a liberal addition of pepper to the brandy-vinegar solution.

Sour Cream

To 20% cream add 2 to 3% skim milk powder. Heat slowly to 120° F. to dissolve the powder and follow this by pasteurizing at 145° F. for 30 minutes. Cool to 70 to 72° F., and add 3 to 5% of

good starter, thoroughly broken up. Dilute 20 drops of commercial renuet extract in about 1/4 glass of water and add this to 100 lb. of cream, agitating it thoroughly to distribute the rennet. rennet helps to form a thick curd and the cream may curdle in a relatively short period. However, you should hold it over night at the ripening temperature of 70 to 72° F, to develop the desired acid flavor. Follow this by breaking up the curd while cooling to 40° F, and hold at this low storage temperature,

Infants' Milk, Synthetic

Sugar	40	g.
Soya Bean Powder	125	g.
Lactose	30	ĸ.
Pennut Oil	20	
Dextrin	20	g.
Egg Yolk, Liquid	50	g.
Calcium Lactate	6	g.
Salt	2	g.
Stir in water before use.		

Soya Bean Vegetable Milk

If the dried beans, preferably yellowseeded varieties, are soaked for a few hours, then finely crushed and boiled for about 30 minutes in the proportion of 3 parts of water to 1 part of mash, a milky emulsion is obtained which is very similar in appearance and properties to animal milk. This liquid, separated out by means of a very fine sieve or cloth strainer, is the Soya Bean or vegetable milk used so extensively in China. Soya bean meal after the oil is extracted or whole soya bean meal may be utilized quite as well as the whole bean. In the absence of animal milk, sova bean milk is used extensively in the fresh state and as the basis of various kinds of vegetable cheeses in oriental countries. Soya bean milk in the form of a powder is a commercial product in some European countries, and in parts of the United States it has been used in special feeding cases. The milk can be used successfully in numerous preparations, such as breads and cakes, in creaming vegetables, in nulk chocolate, and in custards.

After separating the hquid from the solid material, the residue is still very rich in nutritive substances and can be dried and used for cattle feed or made into flour for human food.

Sova Bean Curd

The addition of magnesium or calcium salts or of rennet or lactic acid to soya bean milk when hot precipitates some of

the protein, forming a grayish white curd which settles out, leaving a yellowish water liquid. This card, after being drained and pressed, represents bean curd of tofn, which is extensively eaten and forms the basis of numerous fermented, smoked, and dried cheeses in China and Japan. Bean curd is made fresh daily and is a staple article of diet among oriental peoples. In many cities of the United States having a large oriental population fresh bean card may be found in the Chinese and Japanese markets.

Dry Mix for Making Chocolate Milk in Dannes

Cocoa	1.75	lb.
Cune Sugar	7	16.
Agnr, Powdered	0.14	lb.
Vamilia	0.003	lb.
Sult	0.025	lb.

Mrs the above ingredients well and add to each gallon of milk in the pas-tenrizer at 185° F. Agitate and hold for 1/2 hour.

Cocoa Malt Powder

Cocon Powder	23 lb.
Fine Granulated Sugar	70 H.
Malt Powder, Mild Flavor	20 lh.
Skim Milk (Soluble)	14 lb.
Sodom Bicerbonate	2 oz.
Salt	8 oz.
Vamilla	1/2 02.
Vanilla Extract	1/2 02.

Mix ingredients thoroughly and pass through a coarse sieve. This mixture can be packaged in cans, glass containers, or in 1% oz. envelopes for individual

Stable Phocolate Milk U. S. Patent 1,989,758

In carrying out the process of making the milk staich emulsion, the chocolate, sugars (when the latter are used), starch, and the gum may be introduced, as dry substances, into the milk, thoroughly mixed, and the mixture heated to a temperature of 170° to 200° F., or higher if desired- although this is not necessary in place of temperatures approximating 240° F. heretofore recommended, for periods from 20 to 30 minutes, more or less. Preferably, however, a syrup is first made of the chocolate and sugar, and this syrup, together with a preformed mixture, in proper proportions, of the starch and gum, added to the milk

and the final mixture agitated and heated as described.

As a matter of convenience to the beverage manufacturer, and in order to insure correct proportions between starch and gum, the starch and gum may be compounded together and the compound delivered to the beverage manufacturer.

In making the compound the agaragar, for example, is preferably ground dry and screened to the same degree of fineness as the starch and is then thoroughly mixed with the starch in the proportions indicated by the specific examples given below. In such a mixture the agar-agar, although very small m quantity, approximately from 1 to 20 parts of agar to 100 parts of starch, will remain evenly distributed in the starch. It will not sift out. This novel mixture will disperse in the chocolate vehicle much more easily than if the ingredients were introduced into the liquid as separate substances. If the agar is not finely ground it will swell instead of dissolving, particularly at the low temperatures preferably used in compounding, with consequent loss of stabilizing power.

The following examples of typical mixtures, with preferred percentages of the ingredients, will serve to illustrate the character of the present invention. The percentages are by weight.

Formula No. 1

Milk	90.48
Cane Sugar	4.82
Dextrose (Cerclose)	2 41
Cocon (High Grade, Dark)	1.27
Raw Tapioca Starch (Scott	
Test 150)	1
Agar-Agar	0.02

Any suitable sugars may be used in the suspension or in the dry product or the sugar ingredient may be omitted if desired. The amount of the sugar ingredient may be varied to any extent. For any usable quantity the sugar does not add to the viscosity of the beverage. The amount of cocon or chocolate may also be varied. The matter of taste or of economy will govern any increase or decrease. As much as 2.5% of cocoa may be used without changing the percentage of starch or gum. The starch ingredient may be increased to 2 or 3%. Experience goes to show that 1% is near the critical lower limit. More than 2 or 3% gives too high a viscosity and is likely to give a distinct starch taste to the product. The agar-agar may be varied in amount from about 0.01%, but at the upper limit there is a

strong tendency to segregation in jellylike lumps.

No. 2	
Milk	90.78
Cane Sugar	4.06
Cerelose	2.03
Cocoa (Chenper Quality than in No. 1) Raw Corn Starch (Scott	1.673
Test 100)	1.433 0.024
	0.021

The first four items may be varied as indicated in No. 1.

The same quantity of modified corn starch may be used in place of the specified raw corn starch. The amount of corn starch may vary between 1 and 2%. Where raw corn starch is used the lower limit of the gum quantity should not be quite as low as in No. 1.

		No. 3			
Milk					91
Cane 8	Bugar				4.07
Cerelos	e (Corr	1 Sugar)		2.03
Cocoa	•		′		1.676
Wheat	Starch	(Scott	Test	85)	1.2
Gum		•		•	0.024
			-	_	

The variations may be substantially the same as with No. 1.

The time of cooking with the raw corn starch should be ordinarily 25 to 30 minutes; with the modified corn starch 20 to 25 minutes; with the tapioca and wheat starches about 20 minutes.

Chocolate-Flavored Milk

Cocoa 20 lb. Sugar 90 lb. Skim Milk 90 lb.	this improved formula use:	
Skim Milk 90 lb.	roa 20 H	١.
).
The the above summer and some it.	ım Mılk 90 ll).
To the above syrup and 2000 th, (the above syrup add 2000 lb.	of
milk; heat to 143 1/2° F. and hold for 3	heat to 1431/2° F. and hold for	30

To the above symp and 2000 lb, or milk; heat to 143½° F. and hold for 30 minutes. Homogenize the mixture at 2000 to 3000 lb, pressure while hot.

Cool and bottle.

Joor and bottle.

Non-Settling Coo	oa Milk
Cocoa Powder	6 oz.
Sugar	28 oz.
Sodium Alginate	1 oz.
Milk	15 at.

Mix together the eccon, alginate, and sugar. Heat the milk to 160° F., add the dry mixture slowly with constant stirring, for thirty minutes. Cool the hatch to 45° F. and hold for two hours before bottling in sternlized bottles. The eccoa powder can be of any fat percentage from 10 to 25%. The milk can be either whole milk or akim milk, or any

mixture of each. Additional flavoring ingredients such as vanilla, malt, caramel, etc., may be added.

Boiled Cocoa Frosting

Sugar	16 oz.
Salt	1'16 07.
Water	16 oz,
Vanilla Extract	1/1 OZ.
Dairy Butter	8, 0Z.
Cocoa	3 oz.
Corn Starch	1 oz.
Man manus access a	-134 4 . 41

Mix sugar, cocoa, and salt together, then add slowly 8 oz. of boiling water and when all the water has been added bring mix to a boil. Make a pre mix of corn starch and cold water, then add to the above mix and again bring to a boil Continue boiling with low flame, until the fiosting has become thekened which usually requires 3 or 4 minutes. Remove from flame, add butter and vanille extract, beat well, allow to set and cool.

Chocolate Filling

Milk	8	oz.
Sugar	2	oz.
Flour	11/4	07.
Salt	1/1	02.
Whole Eggs	2	θZ,
Vanilla Extract	1/1	07.
Unsweetened Chocolate	1	υZ,
Hout milk to bailing point	0.1.1	CHIE

Unsweetened Chocolate 1 oz.

Heat milk to boiling point, add sugar,
flour, and salt, stirring thoroughly. Cook
for fifteen minutes, add eggs slightly
beaten, cook for 5 minutes longer. Add
flavoring and unsweetened chocolate, and
1 oz. powdered sugar and stir.

Chocolate Mocha Frosting

Powdered Sugar	112	lb.
Hot Coffce	3	oz.
Unsweetened Chocolate	2	07.
Butter	3/4	oz.

Moisten the sugar with coffee, blend the chocolate with dairy butter. Mix the two blends together and beat until smooth.

Chocolate Icing

Unsweetened Chocolate	2	oz.
Water	4	OZ.
Sugar	16	07.
White of Eggs	2	oz.
Vanilla Extract	1/1	g OZ.
Warm water, then add po-	wdered	sugar

warm water, then add powdered sugar, cook to approximately 216° F. until the mix threads well on the end of a spoon. Stir in the well-beaten white of eggs, then add melted chocolate and vanila

and stir thoroughly to proper consistency.

Builed Marshmallow for Topping Formula No. 1

No. 1	
Granulated Sugar	3 lb
Glucose	12 oz.
Water	1 pt.
Boil to 240° F.	•
No. 2	
Egg Whites	1¼ pt. 8 oz.
Granulated Sugar	8 oz.
No. 3	
Water	4 0%.
Powdered Gelatin	1 oz.
Vanilla Extract	1/2 OZ.
XXXX Sugar	4 07.

Method: Set contents of No. 1 into copper kettle, dissolve well together, and place over moderate fire. Set contents of No. 2 m small 12-quart muchine kettle. Warm contents of No. 3 in small

bowl and thoroughly dissolve.
When contents of No. 1 reach 225° F.,

when concents of No. 1 reach 225° F., start machine going with No. 2 on high speed. Also see to it that sides of copper kettle are kept clear of sugar crystals, by wishing sides of kettle with water and brush.

The meringue content of No. 2 should

The incringing content of No. 2 should be ready about the same time that the boiling content of No. 1 reaches the degree of 240° F.

With the meringue ready, and the boiled sugar at 210° F., pour the boiled sugar on to meringue slowly in thin strenn (this is important). Let the machine run on high speed during this operation.

Now add dissolved contents of No. 3 to the mass, and continue whipping on high until a fine bodied smooth meringue is obtained.

Formula No. 2 (Quicker Method)

No. 1	
Egg Whites	1 pt.
Granulated Sugar	1 pt. 1 lb.
XXXX Sugar	8 oz.
Taproca Flour	1/2 oz.
No. 2	
Glucose	4 0%.
Water	4 02.
Gelatin	1/2 OZ.
Vanilla Extract	1/2 oz.

Method: Dissolve contents of No. 1 all together over double boiler and heat to 120° F. Keep contents stirred with wire hand whip. Now set kettle in machine

and, with wire whip attached, beat on high speed. Immediately dissolve contents of No. 2 by warming, until all are dissolved together, then pour into machine and continue whipping until a fine meringue is obtained.

Whipped Cream for Baker's Topping

The cost of whipping cream and the fact that it will not stand up alone for very long makes its use almost prohibitive.

Fortified Whipped Cream

Cold Water	5	qt.
Meringue Powder	6	oz.
Sugar	4	lb.
Salt	1	oz.
Starch	14	oz.
Gelatin	3/4	oz.
Vanilla Extract	1	07.
Heavy Cream	1	qt.
		٠.

In the machine put 1 qt. of water, the meringue powder and 3 lb. of sugar and whip to just peak (not stiff). Put 3 qt. of water, the remaining sugar and the salt into a kettle and bring to a boil. Dissolve the starch and gelatin in the remaining water, add to the boiling mass and stir until it is thick and clear. Blend the two mixtures carefully with a wire whip and put in the refrigerator until needed. When ready to use, put the mixture into a clean bowl and smooth down with a wire benter. Do not beat. Bring the whipping cream up to about three-fourths stiff, pour it over the boiled mixture, and fold together only until the cream is well incorporated and the mass is smooth.

This should make topping enough for 30 to 40 9-inch pies.

Dokor's Doctin Clare

Datact S 1 CC	tin Ciano	
Pectin	1 oz.	
Sugar	814 lb.	
Water or Fruit Juic	e 21/2 qt.	
Phosphoric Acid	21/2 oz.	

Mix the pectin with some of the sugar. Bring the liquid to a boil and add to the sugar-pectm mixture. Take off the fire and stir until the sugar is thoroughly dissolved. When this has been done add the remaining sugar, stirring in the meantime. Allow the liquid to cool, then add acid.

Coat the berries as much as possible and they will not have a chance to "bleed" and thus soak through into the cake itself. If desired the berries may be dipped into the glaze before they are

applied to the cake and the remaining pectin poured over them so they are nicely coated.

Baking Powder

Sodium Acid Pyrophosphate	42 oz.
Sodium Acid Carbonate	30 oz.
Maize (or Rice) Starch	28 oz.

Stable Baking Powders German Patent 599,493

Formula No. 1

Cream of Tartar	44 g.
Tartarie Acıd	6 g.
Sodium Bicarbonate	27 g.
Wheat Flour	20 g.
Carbamide	1.5 g.
Magnesium Peroxide	1.5 g.
No. 2	
Calcium Biphosphate	34 g.
Sadum Riearbanata	93 %

Magnesium Peroxide

1.5 g.

1.5 g.

Wheat Starch Powder

Carbanide

No. 3	
Sodium Acid Pyrophosphate	44 g.
Sodium Bicarbonate	32 g.
Maize Starch Powder	22 g.
Carbamide	1 g.
Magnesum Peroxide	1 g.
15 g. of above baking poused for 500 g. flour.	owders are

Sova Bean Flour Bread

71 1 1/ 4

Formula No.	1
Soya Flour	65 lb.
Wheat Flour	260 lb.
Sugar	10 lb.
Salt	5 lb.
Yeast	15 lb.
Shortening	15 lb.
Water (Variable)	210 lb.
Mix 3 minutes, ferment	at 90° F.
First punch	45 min.
To bench	15 min.
Proof	45 min.
Bake	30 min.
Temperature of Oven	445° F.
No. 2	
Whole Soya Flour	25 lb.

~	10.
25	lb.
50	lb.
3	lb.
1.75	lb.
2	lb.
2	lb.
	50 3 1.75 2

Sugar	1.5	lb.
Dry Malt	1.5	lb.
Water	about 10	gal.

The straight dough method is employed. A rather wide range in the quantity of water to be used is permitted. This is done to allow for the particular water absorption of the whole wheat flour and the clear that may be used by the laker. A straight dough is made but the whole soya flour is scaked for half an hour with a portion of the water before the dough is made.

"Non"-Staling Bread U. S. Patent 2,009,140

One-half to one per cent arabnose (based on flour) is added to dough.

Infant's Cereal British Patent 416,119

Wheatmeal	52 3	5 lb.
Ontmeal	18	lb.
Cornmeal	10	Th.
Wheat Germ	15	llı.
Salt	0.3	5 lb.
Lucerne	1	16.
Dried Yeast	1	lb.
Bone Meal, Edible	2	lb.
100 ll C l		and

100 lb. of above mixture is cooked with 35 gal, water and then dried on a heated druin.

Storage of Grain and Cereals, Improved British Patent 429,920

1000 lb, of solid carbon dioxide is used per 214 long tons of gram. Both are fed in simultaneously when loading ships or silos.

Chocolate Fudge

Unsweetened Chocolate	6	OZ
Sugar	2	lb.
Milk	1	lb.
Dary Butter	14	oz.
Vanilla Extract	1,10	07.
O. I. I. In the moltad	cho	or.late

Cook slowly the melted chocolate, sugar, milk, and butter mixture to approximately 235° F. until a soft ball is formed when dropped into water. Remove from fire, add vanilla, beat thoroughly until the mass thickens, and then pour into well buttered tin.

Chocolate Cream Fudge

Sugar	11/2	lb.
Corn Syrup	2	OZ.

Unsweetened Chocolate	3	oz.
Salt	1,10	οZ.
Evaporated Milk	8	07.

Heat to a boil, approximately 240° F, the mixed ingredients, until a soft ball is formed when dropped in cold water. Cool to approximately 100° F, and bent to a creamy consistency.

French Candy Bulls

Unsweetened Chocolate	16	07.
Powdered Sugar	2	oz.
Condensed Milk	16	07.
Chocolate Topping	2	αz,

Melt chocolate in double boiler, add signt and stir to prevent lumps. Add condensed milk and stir until smooth, Let set in cool place for two hours. Boll nixture into balls of desired size, and then roll balls in plate of chocolate topping. Let stand over night.

Jellied Fruit Candies

beined range andrea		
Plum Pulp	20	lb.
Peach Pulp		lb.
Cone Sugar		lb.
Corn Syrup		lb.
Powdered Pectin		lb.
Water	2	gnl.

The pectra is mixed well with 5 lb, of sugar. This instance is then starred into the two gal, of water. Cook this solution slowly, to almost the boiling point, with stirring. Then to this smooth solution add the other ingredients. The entire batch is now cooked to 223° F, with stirring. The hot batch may now be deposited in starch molds, and allowed to become cold and firm.

Jellied Orange Candy

Pulp from 50 Oranges.	
Cane Sugar	35 lb.
Corn Syrup	25 lb.
Powdered Pectin	22 oz.
Citric Acid	6 oz.
m: 1	

The pulp is prepared by chopping up the oranges, and then cooked with twee its volume of water until soft, and then rubbed through a coarse strainer, to remove the seeds.

The powdered peetin should be previously mixed with about 10 lb. of sugar. The batch is now cooked to 223° F. Now dissolve the citric acid in a pint of water, add to the batch and once more cook to 223° F. The hot batch is now deposited in starch molds. Allow to become cold and firm.

Jellied Grape Juice Candy

Concord Grape Juice 3 gal.
Cane Sugar 18 lb.
Glucose or Invert Syrup 18 lb.
Powdered Pectin 13 oz.

The pectin is mixed well with about 5 lb. of cane sugar. This mixture is then stirred slowly into the grape juice. The batch is now slowly brought to a boil and then the balance of ingredients are added. Cook to 223° F, with stirring. The hot batch is now run into molds and allowed to cool.

Jellied Apple Juice Candy Apple Juice (from Cooked

Apples) 3 gal.
Cane Sugar 18 lb.
Glucose or Invert Syrup 18 lb.
Powdered Pectin 10 oz.

Citric Acid 5 oz.

Proceed as directed under Jellied Grape Juice Candy.

Jellied Pincapple Juice Candy
This juice can be used in the same
manner as outlined for grape juice.

Candied Sliced Orange, Lemon, and Grapefruit Peels

Slice peel about ¼ in. wide and 3 in. long. Cook peel with several changes of water to remove the bitterness and to make the peel tender. Now add to the peel about a 40% solution of sugar synthetic and cook until the temperature on the thermometer registers 217° F. Now drain the peel and allow to dry over night. The peel may be rolled in granulated sugar if desired. The peel can also be colored red or green with certified food color, if desired. Do this when cooking the peel in the last wash water.

Ginger, Preserved

Drain the ginger well and then cut it up. Place in cold water in a steam-pan, gently bringing to the boil and simmering for twenty minutes. Place in sieves to drain. Transfer to a cold syrup of 4 lb. sugar to each gallon of water, and allow to stand until next day. Transfer all to steam-pan, gently bring to the boil and simmer for 15 minutes. Then place in a clean dry tub and allow to stand until next day. Run off the syrup into the steam-pan and add 3 lb. sugar to each gallon of syrup. Stir well and

bring to the boil. Return this syrup to the ginger in the tub and allow to stand until the following day, then placing in sieves to dry. Roll in sugar and shake out the loose sugar through a coarse sieve. Lastly, spread out to dry.

Preserving Fruit Peels U. S. Patent 1,980,013

A process for treating the rind and peel of citrus fruits comprises heating the material in a sugar syrup for a period not to exceed about 1 hour at a temperature from about 212° F. to about 220° F., placing the material in containers with a relatively small quantity of sugar syrup, heating the containers, while they are unscaled, for a period of about half an hour at a temperature from about 212° F. to about 240° F., scaling the containers, and heating them for a period of about half an hour at a temperature of about 212° F. to about 240° F.

Preserving Red Raspherries by Freezing The best result is obtained by freezing at --18° F. in 50% syrup in either airtight or non-airtight containers, and then storing at --12° F.

Pickling Vinegar Essence

The following is a formula for a concentrated liquid for making pickling vinegar:

Oil of Pimento
Oil of Nutmeg
Oil of Clove
Silve of Capsicum
Acetic Acid (B.P.)
One transport of the street of the

One teaspoonful of this essence is mixed with each quart of vinegar to spice it.

Chewing Gum Bases

a. Bubble Gum Base:

The mixed come and way	
Candelilla Wax	10 lb.
Washed Gutta Soh	75 lb.
Washed Gutta Katian	400 lb.
Washed Pontianac Gum	425 lb.
a Dubbic Guin Dube.	

The mixed gums and wax are heated until the total batch contains only 8-9% moisture.

b. Stick Gum Base:

Pontianac Gum	425	lb.
Gutta Katian	400	lb.
Gutta Soh	75	lb.
Candelilla Wax	60	lb.

Chewing Gum

Formula No. 1

Ball Gum:		
Base b (above)	22	lb.
Corn Syrup	48	lb.
Sugar	117	lb.
Chicle	3	lb.
Wax	11%	
Caramel Paste	21/2	
Flavor	2%	
No. 2		
Penny Stick Gum:		
Base a (above)	40	lb.
Corn Syrup	40	lb.
Sugar	140	lb.
Flavor	30	oz.
No. 3		
Bubble Gum:		
Base a (above)	35	lb.
Pontianac Gum	5	lb.
Corn Syrup	45	lb.
Sugar	115	lb.
Flavor	28	0 Z .

Maraschino Type Cherries

Lambert, Royal Anne, Black Republi-can and Waterhouse varieties can be used. The fresh fruit should show a content of soluble solids in the juice of 16-18% at 21° C. and should be underripe rather than overripe. The bleach solution consists of sulphur dioxide (1.5%) together with sufficient airslaked lime (5.4 lb. per 100 gal. of bleach) to keep the fruit firm and turgid. The cherries lose 7% in weight during the bleaching process. Approximately 250 lb. of cherries is stored in standard 52 gal. barrels and the strength of the bleach solution checked every few days by titration with standard 0.1 N 1 solution. Following bleaching, the cherries are stemmed, graded and pitted. The sulphur dioxide-lime solution is removed by leaching with hot and then with cold water. The sulphur dioxide remaining in the cherries should be less than 0.035%. The dye used for coloring the cherries is No. 80 Ponceau 3R. A solution of % oz. of dye powder in 8 gal. of water is sufficient to color 100 lb. of pitted cherries at a temperature of 93° C. After coloring, the cherries are preserved by gradually increasing the concentration of sugar until a 50% syrup is reached. For flavoring, oil of bitter almonds and amyl acetate are used as desired. Pasteurization of the bottled cherries is effected by a heat treatment of 35 minutes at 91° C. for No. 10 cans holding somewhat less than 1 gal.

Preventing Browning of Peaches After Lye Peeling

Dip in 14% hydrochloric acid for 14 to 1 minute and wash with water.

Non-"Bleeding" Jellies U. S. Patent 1,913,576

To prevent watering of jellies made with pectin, or agar, use 1/2 to 1% sodium alginate.

Jam and Jelly from Fruit Juices

Although most fruits contain small quantities of pectin and acid, many fruits do not contain sufficient amounts of these essential elements to produce jellies when the fruit juices are contained with sugar. A small quantity of malic acid is found in the apple, and a little tartaric acid in the grape. Citric acid is contained in the lemon, the orange, and many other fruits.

Manufacturers of jellies can make high grade pure fruit jelly from all fruit juices by adding a very small amount of fruit acid (either citric, tartaric, or malic), less than one-half of 1%. The addition of small quantities of fruit acid and fruit pectin to fruits which are naturally deficient in these important constituents will improve the fruit flavor in the finished fancy preserve and the standard jum.

There are a few fruits which naturally contain enough acid and pectin to make jellies when the boiling with sugar is continued for 15 or 20 minutes. This excressive boiling, however, evaporates a large quantity of the fruit juice and flavor which should be retained in the finished product. For making jellies from these fruits deficient in pectin and acid, additional quantities of these substances must be added.

Purified powdered pectin is now being made from apples, lemons, and oranges by several firms. The product is very carefully standardized on the beass of jell strength, so that ½ oz. of purified powdered pectin will jell 50 oz. of cane sugar when mixed thoroughly with the sugar and then placed in a suitable cooking pan containing 2½ pints of water. Heat with constant stirring over a strong flame until the mixture weighs exactly 5 lb., then add ½ of a fluid oz. of a 50% solution of fruit said. Mix thoroughly and pour into jelly glasses, Purified powdered pectin of such strength is designated "No. 100."

Pectin syrup is made by mixing thoroughly 5 lb. of powdered No. 100 pectin

with 20 lb. of cane sugar. Place the mixture in a suitable container and add sufficient boiling water to make 10 gal. when the temperature of the syrup is reduced to 70° F. Agitate a few minutes to dissolve the pectin. A 50% solution of fruit acid is made by placing 20 lb. of crystal, granular, or powdered tartaric, or citric acid in a 5-gal. stone jar and adding sufficient boiling water to make 5 gal. when the temperature of the liquid is reduced to 70° F. Agitate the hot liquid until the tartaric acid is dissolved.

All fruit juices for jelly production should have as little added water as is consistent with the proper extraction of pectin, color, and flavor from the fruit being used. Soft juice fruits, such as grapes, require very little, if any, additional moisture. Hard fibrous fruits, such as quinces, require the addition of a relatively large amount of water. In the following formulas for jellies, actual fruit juice is specified exclusive of added water. If water is added to the fruit during cooking, the amount of juice used in the formula should be increased by an amount equal to the quantity of water added at the time the fruit was heated in preparing it for the press, less the small quantity lost in evaporation.

Loganberry, Guava, or Pomegranate Juice Jelly

Filtered Fruit Juice (About 12 gal.) 100 lb. Cane Sugar 97 lb.

or Fruit Juice from 2x1 Cold

Pack Fruit (About 17 gal.) 167 lb. 2x1 cold pack fruit means 2 parts of fruit and 1 part sugar, usually placed in barrels and frozen.

Cane Sugar 30 lb. Fruit Peetin Syrup 1½ gal. 50% Solution Fruit Acid 10 fl. oz.

Crab-Apple, Current, Gooseberry, or Quince Juice Jelly

Filtered Fruit Juice (About 12 gal.) 100 lb. Cane Sugar 97½ lb. Fruit Pectin Syrup 1½ gal. 50% Solution Fruit Acid 12 fl. oz

Cherry, Elderberry, Strawberry, Pineapple, or Raspberry Juice Jelly Filtered Fruit Juice (About 12 gal.) 100 lb. Cane Sugar 96 lb. Fruit Juice from 2x1 Cold
Pack Fruit (About 17
gal.) 167 lb.
Sugar 29 lb.
Fruit Pectin Syrup 2 lb.
50% Fruit Acid Solution 26 ft. oz.

Blackberry, Grape, or Plum Juice Jelly
Filtered Fruit Juice
(About 12 gal.) 100 lb.
Cane Sugar 97 lb.

or Filtered Fruit Juice from 2x1 Cold Pack Fruit (About 17 gal.) 167 lb. Cane Sugar 30 lb. 50% Fruit Acid Solution 15 ff. oz.

Apricot, Peach, or Nectarine Juice Jelly Filtered Fruit Juice (About 12 gal.) 100 lb. Cane Sugar 96 lb. Fruit Pectin Syrup 2 gal. 50% Fruit Acid Solution 23½ fl. oz.

In each formula for fruit jelly, cook the fruit juice, sugar, and fruit pectin syrap to 220° F. at or near sea level, or 8° above the boiling point of water in your factory. Then add the fruit acid solution and mix thoroughly. Fill the jelly quickly into glass and seal at once. If the temperature falls below 18° F, when the container is closed, it should be pasteurized at 180° F. for 20 minutes, if the glass does not contain more than 8 oz. The yield is approximately 164 lb, of finished jelly at 65% soluble solids.

| Standard Cherry Preserves and Jam | Fruit | 82 | 1b. | Cane Sugar | 90 | 1b. | Fruit Peetin Syrup | 2 | gal. | Fruit Acid Solution (50%) | 13½ fl. oz.

In making fancy and standard preserves and jams, cook the fruit, sugar and pectin syrup to 221° F. at or near sea level, or 9° above the boiling point of water at your factory. Then add the fruit acid solution and mix thoroughly. The yield is approximately 158 lb. at 68% soluble solids.

In making standard preserves and jams, cook the fruit, sugar and fruit pectin syrup to 222° F. at or near sea level, or 10° F. above the boiling point of water at your factory. Then add the fruit acid solution and mix thoroughly. The yield is approximately 158 lb. at 69% soluble solids. Fancy preserves jams, and standard preserves and jems

should pass through a cooling pan to re-
duce the temperature to 180° F. and then
duco the temperature and the same
run quickly into glass, and be sealed at
once. Then pasteurize glass containing
once. Then pasteurize glass containing
from 1 to 21/2 lb. at 190° F. for 25
Irom 1 to 279 to. at 100 1. 101 20
minutes. The temperature is reduced to
minutes. The temperature is reduced to
180° F. before running preserves into
100 T. Deroie imming himming
glass so as to prevent the fruit from
floating.

The thermometer should be accurate and should be tested at least once weekly when used daily. A very accurate determination for soluble solids contained in fruit products can be obtained by using a refractometer.

Quince, Damson Plum, Gooseberry or Loganberry Jam

Hoganoenj	o am	
Fruit	100	lb.
Cane Sugar	981/2	lb.
or		
Cold Pack Fruit	150	lb.
Cane Sugar	481/2	Ъ.
Fruit Pectin Syrup	3	qt.
Fruit Acid Solution		•
(50%)	714	fl. oz.
(-··//		

Blackberry, Grape, Strawberry, Raspberry, Pineapple, or Plum Jam (Except Damson Plum) Fruit 100 lb.

 Cane Sugar
 98 lb.

 or
 or

 Cold Pack Fruit, 2x1
 150 lb.

 Cane Sugar
 48 lb.

 Fruit Pectin Syrup
 1 gal.

 Fruit Acid Solution (50%)
 14 d. oz.

 Apricot, Peach, Nectarine or Pear Jam

 Fruit
 100
 lb.

 Cane Sugar
 97½
 lb.

 Fruit Pectin Syrup
 1¼
 gal.

 Fruit Acid Solution
 18½
 fl. oz.

Cherry Preserves and Jam

Fruit 100 lb.
Cane Sugar 90 lb.
Fruit Pectin Syrup 2 gal.
Fruit Acid Solution (50%) 14 ff. oz.

Damson Plum, Gooseherry or Loganberry Jam

y am		
Fruit	82 100	lb.
Cane Sugar or		
Cold Pack Fruit, 2x1	123 571/4	lb. lb.
Cane Sugar Fruit Pectin Syrup	3	qt.
Fruit Acid Solution (50%)	7	fl. oz

Blackberry, Grape, Strawberry, Pineapple, Raspberry or Plum Jam (Except Damson Plum)

Fruit 82 lb.
Cane Sugar or
Cold Pack Fruit, 2x1 123 lb.
Cane Sugar
Fruit Pectin Syrup
Fruit Acid Solution (50%) 13½ fl. oz.

Apricot, Peach, Necturine or Pear Jam
Fruit 82 lb.
Sugar 97½ lb.
Fruit Pectin Syrup 1½ lb.
Fruit Acid Solution (50%) 13½ fl. oz.

Cherry Pie Filler

Put the cherries, sugar, syrup and benzeate into a steam kettle with 7 gal. of water. Bring to the boiling point and then add slowly while stirring, the paste made by mixing the corn starch and tapicca flour with the other 2 gal. of water. Heat and stir until the requisite consistency is attained.

Honey Jelly

 Strained Honey
 24 lb.

 Catrus Pectin No. 80
 4 oz.

 Citric Acid Solution (50%)
 1 oz.

 Water
 1 gal.

Heat the honey to 155° F. in a steam-jacketed kettle.

narketed actue.
In another kettle, heat the water to
180° F. Take about a pint of the honey
and mix it with the dry pectin to make
a smooth paste. Scrape this paste carefully into the hot water and bring to the
boiling point, stirring until all is dissolved.

Add this solution to the honey and mix well. The resulting temperature should be 170° F. If not, raise to this point. Stir in the acid at once and run into containers.

For large size containers, 30 lb. pails or larger, use 20% more pectin.

Plum Jam		
Fresh Fruit	27	lb.
Water	121/2	
Sugar	50	lb.
Pectin	4	0 Z.
Tartaric Acid	11/4	oz.

Sugarless Marmalade for Diabetics

Lemons 144, the peel of one large orange, saccharin 5 gr., water 7 oz., gelatin 46 oz. Wash the orange and lemons, finely shave the skin (avoiding white pith) and chop up small; add the juice and pulp of the lemons. Put into saucepan and cover with the water. Bring to boiling point and simmer for two hours, adding water when necessary to keep to stated amount. Cut the gelatin into fine strips; add it with the saccharin to the mixture and stir for ten minutes. Put into a jar an leave it to set. The keeping properties of this marmalade are not very good, and if it be desired to store it for any length of time a small quantity of sulphurous acid—forty parts per millimen—preferably in form of potassium metabisulphite should be added.

Apple Chutney

Apple chutney is prepared from the fresh apples, peeled, cored, and cut into pieces about half an inch cube. The exact shape of the pieces is not important so long as they are not too small. The apples, after chopping, are allowed to stand over night and then drained from any juice that may have separated, the latter being reserved.

the latter being reserved.

To every 60 lb. of apples 100 lb. of sugar is weighed out, made into a syrup with water, and boiled to 240° F. Into this syrup the small quantity of juice that may have separated is incorporated. While still boiling hot the syrup is poured on to the chopped apples in a suitable container, stirred and allowed to stand for 24 hours. The syrup and apple is then placed in a pan and boiled gently, together with chopped raisins, chopped astem ginger, and as much spice (such as mace, pimento, and nutmeg) and vinegar as taste demands, and the product bottled hot. Served with cold meatparticularly ham and pork—and similar dishes, this chutney is delightful. The color should be golden brown, but this can be darkened if desired with a little sugar caramel. The only machinery required, apart from the boiling pan, is a chopping for dieing machine.

Apple Sauce

Apple sauce, well known in every home as the correct adjunct for roast pork and duck, and usually consisting of apples sliced and stewed with a little sugar, can be truly called a sauce if prepared as follows:

Fresh apples, as green and fresh as can be obtained, are placed in a clean barrel. A steam coil is inserted and the apples cooked for 15 minutes by contact with live steam at about 60 lb. pressure. Care should be taken to see that the steam line is drained before the valve is opened, otherwise the condensed water will enter the barrel and materially affect the consistency of the finished product.

When cooked, the apples are passed through a pulping machine, using the finest sieve obtainable.

To 80 lb. of this apple pulp in a boiling pan add 80 lb. of sugar and 5½ lb. of 80% acetic acid. Stir and cook for 15 minutes. Spices (such as cinnamos, cloves, mace, and a trace of onion or garlie) may be added to suit individual taste, and the product filled into widemouthed bottles.

Apple Butter

Apple butter, which enjoys considerable popularity, is a preparation of a different type, being intended as a spread for sandwiches and at the teatable, and being in fact a kind of concentrated jam.

Processes vary, but consist in the main in expressing the juice from 100 lb, of freshly cooked apples and concentrating with 70 lb. of sugar in a boiling pan to 234° F. At this point 50 lb. of apple pulp, prepared as in the foregoing formula for apple sauce, are added, together with cinnamon, clove and mace spicing, and the mass gently cooked to 228° F.

Prevention of Browning of Fruit and Juices

Treatment with a 0.1% solution of thiourea prevents or retards browning of surfaces of cut fruits. Addition of 0.01% thiourea to apple juice prevents darkening.

Chevon Mince Meats

Oncion minor income		
Chopped Chevon (Cooked		
before Chopping)	10	lb.
Brown Sugar	15	lb.
Washed Currants	15	lb.
Molasses	10	lb.
Granulated Sugar	10	lb.

-			_
Seedless Raisins	15	lh.	
Chopped Apples	30	lb.	
Vinegar, Grape Juice, or		40 ^	
Sauterne Wine	T	Tb.	
Strong Coffee (Percolated			
Preferred)	10	lb.	
Jelly (Apple, Currant, Ra	sp-		
berry, or a Mixture)	` 5	lb.	
Citron	5	lb.	
Salt **	1,4	lb.	
Lemons (Use Junce and			
Grated Rind Only)	139	doz.	
Cook slowly for 3 hours,	addir	ng suff	i-

Cook slowly for 3 hours, adding sufficient water to prevent burning. When cool, add

Rose Water	4 oz.
Cloves	4 oz.
Cinnamon	8 oz.
Nutmeg	4 oz.

Chevon from 8 months to a year old is best for this formula. In using this formula in pies place butter freely over surface before placing top crust.

Salted Soya Beans

A product similar to salted peanuts is obtained as follows:

Soak beans in salted water for 18 hours. Cook beans in lard or vegetable shortening at 170° C, until all water has been driven off and the beans float in the oil.

Fruit Gelatin Powder

 Sucrose
 30 lb.

 Dextrose (Corn Sugar)
 30 lb.

 Gelatin (175 Bloom)
 1 5 - 2 lb.

 Citric or Tartaric Acid
 .75-1 lb.

 Fruit Juice, Fruit and Water 37 lb.
 Flavor and Color

 as desired
 as desired

The gelatin is mixed in the water and dissolved in the usual manner, the sugars are dissolved and at 145° F. mixed with the gelatin solution. Cool to 100° F. and add remaining liquids such as flavoring. color and acid. Let mixture thicken before adding fruits.

Pour into shallow pans to a depth of 1/4 to 1/2 in, and set in cooler. When set turn out and cut into squares.

About 15% by weight of these cubes are stirred into ice cream as it comes from the freezer. The cubes may be added to the ice cream just before withdrawing but some naturally will be broken up.

A slab of the gelatin can be used as a layer in parfait ice cream and the cubes can be used as fillers in fancy pies, etc.

Jelly "Crystals" Formula No. 1

90 1Ъ.
20 lb.
32 oz.
68 OB.
· Aug
81 lb.
5 lb.
4 08.
2 02.
as desired

Gelatinless Jelly Powder

U. S. Patent 1,1	774,474
Agar-Agar, Powdered	1/10 OE.
Sugar	1/16 OZ. 1% OZ.
Tartaric Acid	1/18 oz.
Sodium Bicarbonate	3 gr.
The above forms a stiff	jelly with 8 oz
water.	• •

Lemon Gelatin Powder

	ascaron (iciarii		
Suga	r	10	lb.
Gela	ti n	1	lb.
Citri	c Acid	2.8	OZ.
Leme	on Oil, U.S.P.	11/4	dr.
Certi	fied Yellow Foo	d	
Co	lor	6	gr.
Wate	er	614	gr. fl. dr.

Blancmange Powder

Cornflour	100 lb.
Arrowroot	12 lb.
Color	12 dr.
Flayor	6 oz.

Custard Powder

Cornflour (St. Vincent)	300	lb.
Arrowroot	20-30	lb.
Vanilla	6	oz.
Essence Nutmeg	11/2	dr.
Color Powder	35	dr.

This to be used at the rate of 1½ oz, per pint of milk. The smoothness of the product is increased by the amount of conflour used.

Compound Maple Table Syrups

Cane Sugar—Maple Sugar Blends
Sugar Syrup
Vermont Maple Syrup
Corn Syrup—Cane Sugar Blend

Corn Syrup (39° Bé.) 50 pt. Sugar Syrup 50 pt.

Caramel color to suit.

162	тне снеміса	L FORMULARY	
Cane Sugar—Invert Invert Syrup Sugar Syrup	50 pt. 50 pt.	Glycerin Water Leohol	52 oz. 5 pt. 3 pt.
Caramel color to suit.			nond Flavor
Sugar Syrup New Octobs Molasses	50 pt.	Caramel Color Glycerin, C.P. Benzaldehyde Alcohol	2 0%. 2 0%. 1/2 0%.
Sugar Cane Tab Sugar Lemon Juice	le Syrup 7 lb. 3 oz.	Water Cream S	oda Flavor
Cream of Tarter Caramel Color Sugar Cane Syrup	2 4 g. 5 oz. 4½ pt.	Kanillin Coumarin Alcohol or Glycop Glycerin	5 02 .
Benzoate of Soda Dissolve the sugar in		Water	14 gal.

Dissolve the sugar in bolling water, then stir in the lemon juice and cream of tartar and color; then add the syrup and benzoate of soda. Boil for a few min-utes and strain through fine muslin.

Chocolate Sauce

Unsweetened Chocole	ite 2 oz.	•
Dairy Butter	** 84 OZ.	
Water	4 0z.	۰
Sugar	1b.	•
Vanilla Extract	1/8 oz.	

Melt the crossdate, and add the butter, stir until thoroughly mixed. Then add boiling water gradually with constant stirring "Heat to 280° T. and discontinue heating when a small portion cooled on a dish shows the proper consistency. Cool to approximately 100° F., and add the vanilla flavoring, stir thoroughly.

Apricot	Flavor		
Linalyl Formate		11/2	07.
Amyl Valerianate	•	1/2	oz.
Oenanthic Ether		%	oz.
Aldehyde C ₁₄		1/4	0Z.
Benzaldchyde		1/4	()Z.,
Peach Flavor		8	oz.
Alcohol		1	pt.
Alcohol		67	OZ.
Water		34	oz.

15

Banana Flavoi	ſ	
Amyl Acetate	3	oz.
Butyl, Butyrate	1/8	0Z.
Isobutyi Ketone	1/4	oz.
Ethal Bensoate	1/8	0Z.
Orange Off	1/4	oz.
Benzyl Valerianate	1/6	0Z.
Cinnamon Oil, Ceylon	15	min.
Mace Oil	30	min.
Heliotropin	1/4	oz.

Kola	Beverage	Flavor	
Grain Alcoho		51/2	gal.

Grain Alcohol
Best Vanulla Extract
Oil of Lemon
Oil of Sweet Orango
Oil of Cassia
Oil of Limes
Oil of Nutneg
Oil of Noroli
Extract of Coca Leaves 0Z. 14 7 oz. oz. fl. dr. oz. fl. dr. 3 fl. dr. fl. dr. 1

Allow to stand a month or more and then filter.

Maple Flavor Formula No. 1

rormum No. 1	
Tineture of Foenugreek	6 pt.
Vanillin	% oz.
Musk	1/2 oz.
Balsam Peru	1 oz,
Oil Chamomile	1/2 dr.
Oil Celery	1/2 dr.
Tincture of Coffee	2 pt.
No. 2	
Foenugreek Oleoresin	5 lb.
Hot Water	3 gal.
Alcohol	1 pt.
Malic Acid	15 oz.
Compound Vanilla Extract	10 oz.
Caramel Color	5 pt.
Simple Syrup	150 oz.

Rye Bread Flavor

Cumin Seed, Ground	11 lb.
Anise Seed, Ground	22 lb.
Coriander Seed, Ground	22 lb.
Caraway Seed, Ground	45 lb.

If a liquid flavor is desired the above is percolated with alcohol or if a non-alcoholic flavor is wanted Glycopon S is used.

POOD P	minocia, B	EVERAGES, FLAVORS	163
"Cloudy" Orange Syrup	Concentrate	Housekall E	
Gum Arabic	24 oz.	Household Extracts	
Oil Orange Californian	34 02	(Alcohol	
Oil Lemon Californian	2 300	Pure Lemon	Extract
Orange Color Solution	18 6	Lemon Oil	6.4 Oz.
Simple Syrup	72 oz.	Alcohol, Pure	115 oz.
ated Castor Oil	4 oz.	Water	to 1 gal.
to m	ake 128 oz.	· Pure Orange	Futuret .
Past through colloid mill	_		1 A.
**		Orange Oil	G. C. DR.
	· · · · · ·	Alcohol, Pure	to 1 gal.
Dried Blackberry Con	centrate 🐉		
Dried Blackberries	4 lb.	Pure Almond	Extract
Alco hol	4 pt	l. Oil. Bitter Almond.	The same of
Water	4 pt;	r.F.P.A.	2.35 os.
* **	, 11	Alcohol, Pure	10 . Aug.
Cherry Concentrate,	Natural	Water	to 1 gal
Charries Dried	0 11	Imitation Vanill	
Cherries, Dried Alcohol	8 lb./	Townsia.	70
Water	4 pt.	Vanillin Coumarin	70 oz.
	4 pt.	Alashal Duny (954)	4∕4 ∪Z.
Put cherries in water, he	rat, cool, and	Alcohol, Pure (25% 1 Volume) Simple Syrup	oy 25 gal.
dd alcohol.		Simple Syrun	25 gal. 80 oz.
		Water and Color	to 100 gal.
Cognac Essence	3		
Cognac Ether		Imitation Lemon	n Extract
Rum Ether	650 g. 650 g.~	Citral	34 oz.
Sweetened "Saltpeter Spin	it'' 165 g.	Alcohol, Pilre	5 pt.
Sweetened "Saltpeter Spir Ethyl Acetate	165 2.	Water	to 1 gal.
Oenanthic Ether	5 g.	• · · - · · · · · · · · · · · · · · · · 	
Sugar Color	335 g.	Company D	4-0-4
Alcohol (90%)	4000 g.	Caraway En	CLINICE.
, , ,	Ü	Formula M	ot 1
**		Oil of Caraway in the	.™ 8 g.
Rum Essence			₹ . 50 g.
Rum Ether	200 g.	Glycerin "	" 6 g.
Ethyl Acetate Cinnamon Tineture	40 g.	Water	41 g.
Cinnamon, Tineture Catechu, Tineture Vanillin, Tineture Ethel Formate	10 g.	No. 2	
Catechu, Tincture	10 g.	Oil of Caraway	3 g.
Vanillin, Tincture	10 g.	Alcohol	80 2
Ethyl Formate	75 g.	Water	20 AFT
Angenca Koot, Tincture	2 g.		-
Angelica Root, Tincture Peruvian Bark, Tincture Orange Flower Water	10 g.	Cardamom E	xtract
Washeng Formus	30 g.	Oil of Cardamom, Co	
Woodruff Essence	30 g.	Alcohol	50 g.
Butyric Ether	20 g. 650 g.	Glycerin	6 g.
Alcohol (90%) Rum	1000 g.	Water	41 g.
want	1000 g.	*** 4101	34 B.
Rock and Rye Whisky		Cassia Ext	ract .
Grain Fusel Oil Rectified	340 g.	Formula N	o. 1
Green Wine Lees Oil	12 g.	Oil of Cassia Rectifie	
Peru Balsam	12 g.	Alcohol	50 g.
Jamaica Rum Essence	12 g.	Glycerin	6 g.
Vanillin	6 g. 1	Water	41 g.
Ethyl Acetate Coumarin	12 g.	11 4101	5.
Coumarin	15 g.	No. 2 (Cinna	months.
Raisin Wine Essence	580 g.		
		3% Stands	
Peach Essence	8 g.	03 4 4 1 2	
Peach Essence	50 g.	Oil of Cassia Cinnamo	
Peach Essence Bitter Orange Extract Cinnamon Oil Clove Oil		Oil of Cassia Cinname Alcohol Water	on 30 g. 200 g. 170 g.

Extract Celery		Extract Juniper	
Formula No. 1		Oil of Juniper	2 g.
Celery Oil	0.6 g.	Alcohol Water	90 g.
Alcohol	600 g.	Water.	8 g.
Water	400 g.	D	
	200 80	Banana Oil (Synthe lb. oz.	
, No. 2		Dameul Asstate 0 15	32 3 54
Oil of Celery	0.5 g.	America Acceptance 4	3 54
Alcohol	60 g.	Benzyl Acetate 2 15 Amyl Acetate 4 4 Heliotropin - 1 Vanillin - 1	2 58
		Henotropin - 1	2 58
Glycerin	δg.	Vanillin - 1	2 58
Water	34 g.	Butyl Laurate 2 8	3 16
		Geranyl Acetate	2 12
Wild Cherry Ext	act	Amyl Acctate 4 4 Heliotropin - 1 Vanillin - 1 Butyl Laurate 2 8 Geranyl Acctate - *Terpeneless Lemon	
Wild Cherry Bark	8 lb.	Oil	1* 16
Alcohol	4 lb.		
Alconol Water	4 lb.	Blackberry Oil	ř
	4 10.	Vanillin	2 g.
Percolate and filter.		Coumarin	3 0
		Heliotropin	2 g.
Ciunnau 12-1	-4	Methyl Salicylate	2 g. 2 g. 1 g.
Cinnamon Extra		Mothel Anthonilet	≟ g.
Oil of Cinnamon, Ceylon	_3 g.	Methyl Anthranilate Orris (10% Solution)	ī g.
Alcohol	50 g.	Offis (10% Solution)	og.
Glycerin	6 g.	Coriander Oil	. 0 g.
Water	41 g.	Fennel Seed Oil	18 g.
	0	1 Amyl Kutyrata	1 g. 5 g. 6 g. 18 g. 112 g.
/M		Ethyl Benzoate Amyl Acctate Ethyl Acctate Aldehyde C ₁₆	200 2.
Clove Extract		Amyl Acetate	192 g.
Formula No. 1		Ethyl Acetate	397 g.
Oil of Cloves	3 g.	Aldehyde C ₁₆	4 g.
Alcohol	50 g.		U
Glycerin	a g.	Brandy Oil	
	6 g.		
Water	41 g.	Green Cognac Oil	20 g.
		Oenanthic Ether	80 g.
No. 2		Rum Ether	80 g.
Clove Oil	20 g.	Fusel Oil	20 g.
Alcohol	650 g.	011 11111 01	_
Water	350 g.	Oil Wild Cherry	
	6'	Formula No. 1	
0-1-1-7		Benzoic Acid	4 g.
Coriander Extra		Benzaldehyde	6 g.
Formula No. 1		Amyl Butyrate Ethyl Acetate	6 or.
Oil of Coriander	3 g.	Ethyl Acetate	24 g.
Alcohol	50 g.	Ethyl Benzoate	24 g.
Glycerin	6 g.	20071 2000000	K.
Water	41 g.	N. 0	
***************************************	-1 R.	No. 2	04
		Amyl Acetate	24 g.
No. 2		Amyl Acetate Amyl Butyrate Ethyl Benzoate Benzaldehyde	12 g.
Oil of Coriander	3 g.	Ethyl Benzoate	12 g.
Alcohol	80 g.	Benzaldehyde	32 0.
Water	20 g.	Oil Sweet Orange Calif.	4 g.
	~ ∨ 8.	Oil Cloves	3 g.
Ginger Ale Extra	sct	Cherry Oil (Synthet	ic)
Oleoresin Capsicum			
Safrol	1 02		dr. mir
Cinnamiat Aldohyda	1 04	Benzylidene For-	
Cinnamic Aldenyde	1 oz. 1 oz. 1½ oz.	mate 1 -	
	1½ oz. 1½ oz.	Oenanthic Ether 4 8	
Mace Out		Ethyl Methyl An-	
Citral	179 02.	Littly Laterny Later	
Citral Geranyl Acetate	1/4 oz.	thranilate 1 6	3 12
Mace Out Citral Geranyl Acetate Alcohol One ounce will flavor five	1/4 oz. 1 pt.	thranilate 1 6 Benzaldehyde, F.F.C. 3 1	3 12 4 48

		100
Oil Cognac		02 P Put
Tincture of Prunes	490	Oil Pear Ethereal
Ethyl Butyrate	480 g.	Benzyl Propionate 1 oz.
Oil Cognac	21. g.	Amyl Acetate, Pure 11 oz.
Oenanthic Ether	28 g. 42 g.	Butyric Ether, Absolute 4 oz.
	an R.	//G
. 011 4.0		"Scotch" Whisky Oil
Oil of Green Cogn	ac	Fusel Oil Rectified 510 g.
Sebacic Ether	5 g.	Cade Oil 84 g.
Pelargonic Ether	2 g.	Ethyl Butyrate 445 g.
Cognac Oil	3 g.	Bitter Almond Oil 20 g.
Oenanthic Ether	90 g.	Sweet Almond Oil 20 g.
		Guaiacum Oil 10 g.
Cola Oil for Beverag	res *	0.7 (1)
Oil Lemon	120 g.	Oil Strawberry (Synthetic)
Oil Sweet Orange	80 g.	oz. dr. min.
Oil Nutmeg	40 g.	Ethyl Acctate 42 5 15
Oil Cinnamon	40 g.	Aldehyde C ₁₆ 23 3 40
O11 (1)	20 g.	Amyl Acetate 12 6 24
Oil Neroli, Artificial	40 g.	Ethyl Butyrate 9 - 27 Amyl Butyrate 9 - 27
Alcohol (75%)	15,360 g.	Amyl Butyrate 9 27 Propyl Iso Butyrate 58 5 15 Ethyl Formate 1 2 13
	, 6.	Propyl Iso Butyrate 58 5 15
G 0'1		Ethyl Formate 1 2 13
Curacao Oil		Oil Cognac, Green - 6 47
Benzaldchyde	15 g.	Phenyl Butyl Ketone 2 1
Oil Cassia	30 g.	011 70 11 11 11 11 11 11
Geraniol Extra	30 g.	Oil Raspberry (Artificial)
Linalyl Acetate	50 g.	lb. oz. dr. min.
Petitgrain Oil	75 g.	Ten Rone, Oil - 9 4 45
Orange Oil	650 g.	Aldehyde C ₁₆ - 11 5 25
Lemon Oil	150 g.	Amyl Cinnamic
-		Formate 1 6 6 10
"Holland" Gin Oi	1	Vanillin - 3 20
Lemon Oil	3 g.	Amyl Acetate - 10 3 25
Anise Oil	3 g.	Ethyl Butyrate - 8 2 44
Angelica Root Oil	16 g.	Ethyl Formate - 8 2 44
Fusel Oil Rectified	12 g.	Ethyl Acetate - 12 4 6
Rosemary Oil	16 g.	Iso Butyl Acctate 2 11 6 20 Iso Cinnamic
Coriander Oil	13 g.	Acetate 1 7 6 11
Juniper Berry Oil	940 g.	Amyl Butyrate - 8 2 44
		Imy Ducyrate - 0 2 44
"Old Tom" Gin Oi	1	Concentrated Foam for Beverages
Coriander Oil	270 g.	Saponin 16 oz.
Anise Oil Rectified	80 g.	
Juniper Berry Oil Rectified	610 g.	Glycerin 64 oz. Distilled Water 64 oz.
Caraway Oil	20 g.	
Angelica Root Oil	15 g.	Use 1 oz. to 15 gal. syrup.
	ū	0 # : 5 0 #
Oil Grape (Synthetic	`	Caffein-Free Coffee
		U. S. Patent 2,023,333
lb. oz. Oil Cognac Green - 14	dr. min. 5 26	Ground raw coffee is extracted with a
	5 26	warm mixture of
Methyl Anthran- ilate 7 2	3 55	gg-dichlorethane and
Ethyl Cinnamate - 7	2 43	αβ-dichlorethane
Propvl Cinnamate - 5	6 58	
	4 55	Artificial Mineral Water
		Austrian Patent 142,032
OHE: IS:		
Oil Kümmel Danzig		1 liter of following solution is mixed
Carvol	300 g.	with 10 liters of carbonated water:
Coriander Oil	3 g.	Salt 0.02 g.
Orange Oil	3 g.	Magnesium Sulphate 0.02 g.

Dihydrogen Sodium Phosphate Potassium Nitrate	0.02 g. 0.008 g.	Essence Orange Sulphurous Acid	5 oz. 4 oz.
Calcium Oxide	0.2 g.	Tonic Water	•
		Quinine Bisulphate	8 gr.
Lime Barley Wat	er	Aerated Lemonade	4 pt.
Syrup (66°)		Aerated Water	4 pt.
Barley Extract	2 gal. 3 qt.		
Refined Lime Juice	1 qt.	Lemonade Cryst	als
Citric Acid Powder	7 oz.	Sugar	100 lb.
or		Lemon Juice Powder	
1-2 Solution	14 oz.	Tartaric Acid	4 lb.
Essence Lime	3 oz.		•
Sulphurous Acid	3 oz.	Orangeade Crys	tals
Lemon Color	½-1 oz.	Sugar	100 lb.
Visit management and the residence		Orange Juice Powder	6 to 8 oz.
Orange Barley Wa	ter	Tartaric Acid	4 lb.
Syrup (66°)	3 gal.		•
Orange Concentrate 6-1	1 pt.	Lime Juice Cry	
Orange Beverage Base	7 pt.	Sugar	100 lb.
Barley Extract	1 gal.	Lime Juice Powder	
Orange Color, if Desired	2-6 oz.	Tartaric Acid	4 lb.

SUGAR TABLE FOR SODA WATERS

Pounds of Suga Added to	Q	uantity of		Sugar Percentage		Degrees Baumé
1 Gal. Water	Syr	up Obtained		in Syrup	Density	at 60° F.
	gal.	pt.	oz.			
1	Ĩ.	-	10	103/4	1.043	6
$\frac{2}{3}$	1	1	4	1914	1.080	11
	1	1	14	261/2	1.113	151/2
4 5	1	2	- 8	32%	1.142	18
5	1	3	2	371/2	1.166	201/2
6	1	3	12	41%	1.188	23
7	1	4	6	45%	1,209	25
8 9	1	5		49	1.227	26%
9	1	5	10	52	1.244	281/4
10	1	6	4	541/2	1.258	291/2
11	1	6	14	57	1.271	30%
12	1	7	8	59	1.284	32
13	2	0	2	51	1.296	33
14	2	0	12	62 34	1.306	33%
15	2	1	6	641/4	1.315	341/2
16	2	2		651/2	1.324	34%
17	2	2	10	671/4	1.332	35 1/4
18	2 2	3	4	681/2	1.340	35%
19	2	3	14	69 %	1.347	36

Aging Alcoholic Liquors
U. S. Patent 1,963,165

About a pound and a quarter of potassium permanganate crystals are dissolved in an appropriate amount of water, for example three and one-half gallons. To this solution there is added to about a pound of sulphuric acid, pref-

comes in charred barrels provided with a removable bung. In operating according to the present process, the bung is removed from the barrel and the aqueous mixture resulting from mixing sulphuric acid and potassium permanganate in so-lution is added to the contents of the barrel. Thereafter the bung is replaced and the barrel and its contents are allowed to mature for a short period of time at an elevated temperature. Ryo and bourbon are allowed to mature for about three days at a temperature of 120° F., while rum and brandy are allowed to mature for about two days at the same temperature. When using a lower temperature, for example 100° F., rye and bourbon can be allowed to mature for a period of five days, and rum and brandy for a period of four days. The important point is that after the treatment of the raw alcohol liquor with the treating solution there should be a short maturing period. The function of the elevated temperature is to accelerate the maturing period, and therefore, if the temperature is reduced, the maturing period at this point becomes longer and tree versa. If the temperature is increased above 120° F., the maturing period can be shortened. Of course, the upper temperature limit cannot be too high, since the treatment mixture at highly elevated temperatures would deteriorate the quality of the alcoholic

When the raw alcoholic liquor is treated with the aqueous solution resulting from mixing sulphuric acid and potassium permanganate, there is immediately set up in the liquor a substantial agitation, acting to eliminate the poisonous components of the fusel oils including the aldehydes and the higher alcohols while leaving the esters of the fusel oil to which the arromatic flavor of the liquor is due substantially unimpaired.

After the treated alcoholic liquor has here allowed to mature, as set forth above, the temporary bung is removed. When the bung is removed from the barrel, the chemical and physical action which the liquor is undergoing is very apparent. Immediately upon removal of the bung, there is an evolution of vapors and gases, these representing partial reaction products of the treatment process up to this point. A portion of the impurities present in the original raw liquor have been removed by virtue of the absorptive capacity of the porous lining of the barrel which, as stated, is also in a charred condition, thus augmenting the initial absorptive capacity

of the porous wood of which the barrel is made.

Immediately upon removing the temporary bung from the barrel there is added to the treated alcoholic liquor an agent which will function to bleach and stop the chemical and physical activity taking place in the liquor which has been treated with the sulphuric acid and the permanganate mixture. While various agents may be used to effect the bleaching and the cessation of chemical and physical activity in the alcoholic liquor, it has been found that most satisfactory results are obtained by the addition of an oxygen evolving agent. While the preferred oxidizing agent is hydrogen peroxide, other compounds which are the chemical equivalents of hydrogen peroxide may be used.

The amount of the bleaching and activity neutralizing agent which is added to the treated alcoholic liquor will of course vary with the character and quality of the initial raw product and with the amount of the sulphuric acid permanganate solution which has been initially added to the raw liquor. When adding the sulphuric acid permanganato treatment agent in the proportions above set forth to about 50 gal. of raw liquor, it has been found that the addition of eight onnees of 30% hydrogen peroxide gives very satisfactory results.

After the addition of the bleaching

and activity neutralizing agent, a permanent bung is inserted into the barrel and the treated alcoholic liquor allowed to further mature, preferably under an The following elevated temperature. maturing procedure has been found to give most satisfactory results. maturing rum and brandy, the barrels of alcoholic liquor treated in accordance with the previous steps of the process are maintained in a warehouse having a temperature of about 120° F. for about three weeks. Thereafter the temperature is reduced to about 100° for another week, and then to about 80° F. for an additional week. The warehouse or room in which the liquor is being matured under elevated temperature is then allowed to cool off to normal temperature which usually takes about a week or ten days, unless artificial means are used for cooling the temperature of the storage room. In general this period of maturity varies from about 6 to 8 weeks, and the resulting rum and brandy has reached full ma-turity, having a flavor and mellowness equivalent to rum and brandy which have been naturally aged for a period of approximately four years.

When rye and bourbon are treated, due to the higher content of impurities including fusel oils present in the raw alcoholic liquor, a longer period of maturing is necessary. When rye and bourbon have been treated as above set forth with the sulphuric acid potassium permanganate solution, and then later on after a short period of maturing treated with the bleaching and activity neutralizing agent, the so treated material is subjected for a period of about two months to a temperature of about 100° F. The temperature of the storage room containing the barrels of treated liquor is then reduced to about 100° F. and the treated liquor allowed to mature for about an additional two months. Thereafter the temperature of the storage room is reduced to 80° F. for a period of one month. The storage room is then allowed to cool off to about 70° F., it taking about one month under average conditions for the storage room to reach this temperature, although it is recognized that the cooling may be accomplished much quicker by artificial cooling means.

Kümmel Danzig		
Carvol	300	g.
Coriander Oil		ğ.
Orange Oil	3	ğ.
Alcohol	5000	g.
Water	2250	ğ.
Glycerin	274	ğ.
		-

Methyl, Isopropyl and Amyl Alcohols, Tests For

0.1 g. of vanillin is disolved in 10 mils of alcohol in a test tube and 1 mil of pure sulphuric acid carefully run down the side of the test tube to form a layer at the bottom. By slightly rotating the tube the alcohol and acid are cautiously mixed (care is needed otherwise the sudden rise in temperature will cause violent ebullition) and the colors formed at the area of contact and of the final mixture are noticed.

Methyl Alcohol:	Area of contact Final mixture	l'ale mauve Pale mauve
Ethyl Alcohol:	Area of contact Final mixture	Lemon yellow Colorless
Isopropyl Alcohol:	Area of contact	Bright red towards the acid layer changing to deep blue towards the alcohol layer
	Final mixture	Prussian blue
Amyl Alcohol:	Area of contact	Dull red towards the acid layer changing to deep blue towards the alcohol layer
	Final mixture	Prussian blue A white precipitate
		also forms

In order to differentiate more accurately between isopropyl alcohol and amyl alcohol 10 mils of water are added to each mixture and then shaken well. With isopropyl alcohol the mixture becomes pale blue but rapidly fades, becoming water-white. With amyl alcohol the mixture separates into two layers, the upper alcoholic one being deep grass green (permanent after two hours) and the lower aqueous layer water-white. The white precipitate settles to the bottom.

Approximate Estimation

For the approximate estimation of methyl, isopropyl and amyl alcohols in ethyl alcohol the quantity of sulphuric acid used was increased to 3 mils. Dilutions of the three alcohols in ethyl alcohol were made, 1 in 10, 1 in 100, 1 in 1000 and 1 in 10,000, also a control of ethyl alcohol alone, the color obtained with the latter and the solution of vanillin being a distinct yellowish green.

	1 in 10	1 in 100	1 in 1000	1 in 10,000
Methyl Alcohol	Blue green	Very faintly blue green	Yellowish green	
Isopropyl Alco- hol	Blue	Pale blue	Blue green	Yellowish green
Amyl Alcohol	Deep blue	Blue	Pale blue	Very faintly blue green

Preserving Brewer's Yeast Yeast will keep well indefinitely if covered by a 10% cane sugar solution.

Seed Yeast for Production of Commercial Yeast U. S. Patent 2,016,791

After mixing about 4 lb. yeast with an aqueous aerated "cream" formed by agitating about 12 oz. of calcium sulphate with water, 0.5 to 1.0% of corn starch is added to the mixture, it is maintained at a temperature of about 28° C. for about 30 hours, then diluted with aerated water and allowed to stand for about 18 hours to produce sporulated and durable yeast.

Isinglass Finings for Beer Clarification British Patent 432,159

Pieces of isinglass are steeped in acidifled water for several hours and then gently stirred continuously for 12-15 hours. The liquid, which then has the consistency of thin treacle, is strained through a fine sieve.

Home Made Wine

To two volumes of water in a large glass bottle, add one volume of washed whole grapes and one volume of sugar. Stopper with a cotton plug, place in a warm place, shake up well daily, and allow to ferment for about 8 weeks or until the evolution of gases ceases. Then siphon off or decant, sweeten to taste, bottle and set aside to age.

Bee Wine

Four ounces of sugar and 4 oz. of treacle are mixed with 11/2 pt. of water to form the mother liquid. Small pieces of the ginger beer plant are then added, and the mixture is kept in a warm place. Each day about a tea-spoonful of sugar is added, there is brisk fermentation and a palatable drink is soon ready. The ferment quickly increases, and can be used to prepare a new batch.

Orange Wine

Cut well ripened oranges in half and squeeze out juice. Strain out coarse pulp and seeds. Add 150 p. p. m. of sulphur dioxide; corresponds to about 2½ and stop the treatment when the de lb. of metabisulphite or about 1½ lb. of sulphur dioxide per 1000 gal. of the first on a small scale, in order to avoid

juice. Mix well. Add sugar to increase the Balling degree to 22-24° Balling for a dry wine of medium alcoholic content and to 32-33° Balling for one that will contain a small amount of sugar after fermentation is complete.

Ferment large quantities in open redwood vats, artificially cooling the fermenting liquid, if necessary, to maintain the temperature below 85° F. Smaller quantities are fermented in oak puncheons or barrels. Take the Balling degree once a day to follow the course of fermentation.

When fermentation becomes slow and is nearing completion transfer from the open vat to a covered redwood tank, leaving the bung hole open. Fit with a fermentation bung in order to give a slight pressure of carbon dioxide gas in the tank and thus prevent the growth of vinegar bacteria. Similarly equip bar-

rels or puncheons.

When gas is no longer given off re-move fermentation bung and fill the tank, puncheon or barrel with fermented orange juice and scal with an ordinary bung. Once or twice a week for several weeks loosen the bung for a few seconds to release accumulated gas pressure until fermentation ceases.

Then let stand for two or three weeks to settle, with bung tightly in place. Next drain off, that is, rack from the sediment; this can be done through a bronze spigot inserted in a bung hole near the bottom of the tank, or by syphoning by hose from smaller container. Transfer to clean cooperage that has been sulphured (in which a sulphur wick has been burned). Fill these containers completely full. Let settle two or three weeks. Then rack. Filter clear, This is easily done, usually by means of a pulp filter. The wine can then be polished brilliantly clear through a porcelain candle, or pad type polishing filter. It should next be aged, in wood as is done with grape wine. If new wood is used the tanks or barrels should be soaked out with dilute soda ash solution and water before use in order that the wine will not acquire too strong a wood taste.

If to be rapidly aged, heat to 120° F. a few days in the presence of about 1/4% by weight of oak chips if in redwood, and a head space of about 10%. If in oak barrels no chips are needed. Pump "grief" and loss by improperly rapid aging of a large quantity.

After aging, the "wine" may need a polishing filtering again. After filtering let it rest in wood a few days to "recover" before bottling.

If a fortified wine is desired a special permit or license is required, numerous regulations must be met and numerous forms filled out, either to install a still and use brandy made on the premises for fortification, or to buy fortifying brandy of high proof. Having conformed to all regulations, etc., then brandy may be added to bring the wine to 20-21% alcohol. "Angelica" type sweet forti-fied orange "cider" should show about 10% sugar by chemical test and sherry type 2-4% sugar by chemical test. The former is aged like dry wine; the latter is heated at 130-140° F. for 2-3 months to acquire a sherry flavor and color. By gentle aeration the time can be greatly shortened.

"Champagne" type sparkling orange wine can be made by fermenting juice of 21-22° Balling dry; filtering; aging a few months; adding 2% of cane sugar; fermenting in the bottle with Champagne yeast, disgorging and refilling the bottles; or by fermenting in bulk by the Charmat or other bulk process; filtering and bottling under carbon dioxide pressure.

Or the orange "wine," sweet or dry, can be carbonated with carbon dioxide gas in one of several types of carbonating machines.

In order that non-carbonated, non-fortified sweet "wine" after bottling will not undergo bacterial spoilage it may be preserved with about 300 p.p.m. of sulphur dioxide, or by pasteurizing in the bottle at 140° F. for 30 minutes.

Berry "Wines"

Here the procedure is somewhat different than in making orange "wine." Use ripe, sound berries, sorting out moldy fruit. Crush into open vats. Add 8 oz. of potassium metabisulphite or 4 oz. of sulphur dioxide per ton, or about 2½ h. of the former or 1½ of the latter to each 1000 gal. The metabisulphite is dissolved in water 8 oz. per gal. before addition. Add to the juice. Mix well. Wait 2 hours. Add a starter or 2-3% pure yeast culture. Stir or punch three times daily until the Balling degree drops to about 1/2 or 1/4 the original Balling degree. Fermentation extracts the color and tannin and softens the fruit.

Press in a rack and cloth press. the juice add for a dry "wine" 15% by weight of sugar; for a sweet "wine" about 25%; that is to 1000 gal. of the juice about 1350 and 2250 lb. of sugar respectively. See that it all dissolves.

Ferment and treat as described for

orange "wine."

Rhubarb Wine

Run 32 lb, rhubarb through a meat chopper, strain the juice into a vat and add 6 gal. water. Let stand for 2 days and strain. Let stand for 1 to 2 days, siphon off the clear liquid into a keg and add 24 lb. sugar. Boil up 2 lb. raisins in a little water and add together with 1 lb. sugar coloring. Add also a little gelatin as clearing agent. Let ferment for about 14 days, or until complete. Fill up keg with water and let stand for 4 months before tapping.

Dehydration of Fresh Soya-Slime German Patent 602,935 and 599,639

Example of a Soya-Mud of composition:

Water 50 oz. 40 oz. Lecithin 10 oz. Soya-Oil Warm Soya-Slime 100 oz.

to 60° C., and add

Glycerin Containing Dry Sugar (until sp. g. = 1.36 to 1.39)

Stir thoroughly 1/4 hour, allow to stand. Two layers formed, the heavy one:

Glycerin + Water + Sugar

the light one:

Lecithin + Oil +

Water

Repeat to get a water-content of 10%.

Defoamer for the Sugar Industry Prevents foaming when "saturating" the lime-containing "thin sap." Woolfat, Neutral.

For the Alcohol Industry:

Coconut Oil 20-15 Vaseline Oil

Preservation of Coffee U. S. Patent 1.956.290

Oxidation and "staling" of coffee is curtailed by addition of 0.3% sodium pyrosulphate.

> Denaturation for Food Salt (per 100 kg.) Formula No. 1

Mineral Oil

No. 2 No. 3

0.25 kg.

Iron Oxide Soap Powder

1 kg.

0.25 kg.

For the Chemical Industry

No. 4

Sodium Sulphate, Crystallized

5 kg.

Sodium Sulphate, Calcined 2.5 kg. No. 5

Sodium Carbonate

No. 6 Crystal Ponceau 6R

2 kg. 0.5 g.

Non-Caking Salt

British Patent 407,829

The addition of up to 7% potassium chloride to granular table salt prevents caking.

Non-Caking Sugar

Caking of sugar is prevented by addition of 1% tricalcium phosphate.

Improving Liquid Honey

Heat honey to 71° C.; cool rapidly to 24° C.; add fine crystallized honey with stirring for 15 minutes; cool and bottle.

Non-Mottling and Non-Hardening Maple Sugar

U. S. Patent 1,970,870

Maple sap or syrup is boiled in an open vessel until the temperature reaches 125° F., then allowed to cool, and continuously stirred until cold. The crystallized mass obtained, containing about 2% of water is pressed into blocks occupying 30-31.4 cu. in. per lb.

Clarifying Cider

Pectin (20-30 oz.) is added to 1 gal. of warm cider and the mixture shaken

at intervals for 20 minutes. The strained liquor is added to 100 gal. of cider to be clarified, and after 15 hours at approximately 21° C, the cider is siphoned off, mixed with 2-3 lb. of diatomacoous earth, and filtered through canvas.

Wax Coating for Citrus Fruit U. S. Patent 1,940,530

Fresh fruit (notably citrus) is improved in appearance and made less liable to wither if a thin film of molten wax is rubbed on to the surface (e.g., 5-15% of carnauba wax in paraffin wax at 77-105° C, rubbed on for 10-30 seconds). Advantageous results are obtained if an alkaline wash has preceded this treatment.

Curing Ripe Olives U. S. Patent 1,928,229

Wash olives in 1/2 to 2% caustic soda solution then in water till neutral. Sonk in 1/2 to 5% pyrogallol for a few hours. Without rinsing sonk in 1% caustic soda solution until skin is penetrated; expose to air until black; wash till free from alkali and then soak in brine to develop flavor.

Storing Walnut Meats

Bleached nuts are preserved by packing in earthenware containers with alternate layers of coconut fiber and a 9 to 1 mixture of salt and sodium diliydrogen phosphate crystals.

Vitamin B Concentrate Japanese Patent 101,137

Rice bran or other similar vegetable material is extracted with methanol at 60° C. The solvent is distilled off in vacuo. The extractive residue contains a good percentage of vitamin B.

Detecting Cold Storage Eggs

By dipping eggs in lamp black, one can tell immediately whether they are freshly laid or cold storage.

The test depends upon the fact that storage eggs are treated with an oil to preserve them. If it is a cold storage egg, the lamp black will cling readily to the outer shell, while the amount of lamp black adhering to a fresh egg is said to be negligible.

	Effects of various factors on the vitamin	Long exposure to air, especially at high temperatures, may result in destruction of vitamin A, but it is not readily destroyed by ordinary cooking or canning processes. The yellow coloring matter, carotin, which is found in carrots and in other yellow and green vegetables and fruits, may be changed to vitamin A in the body. Carotene is less readily destroyed by exposure to air and to high temperatures than is the vitamin A in animal fats Since vitamin A in animal fats Since vitamin A in water), it is not lost in cooking water, as are some of the 'water soluble', vitaminal.	Ordinary cooking and canning processes do not destroy vitamin B readily, but since vitamin B is "water-soluble," amin B is "water-soluble," in the cooking water or vegetable juice is thrown away. The addition of sode in cooking vegetables increases the destruction of vitamin B Dring apparently does not destroy vitamin B
VITAMIN DATA	Good sources	Cod-liver oil, halibut-liver oil, salmon and other fish oils Butter Liver and kidney. Egg yolk Whole milk, cream, and cheese made from whole milk. Carrots, pimento perpers, spinsch and other green leaves, and tomatoce Usually, foods having a yellow or green color. Thus green leaves, yellow corn, and sweet potatices are blanched leaves, white corn, and white potatices.	Whole grains Dried peas and beans Nuts Green leafy vegetables Tomatoes Milk Liver Egg yolk
VITAM	Functions in the body	It is essential for: Growth Good health at all ages Successful reproduction Maintenance of healthy membranes which provide a barrier against the in- vasion of bacteria Its absence causes: The surface coverig in various parts of the body to brank down. This may al- low bacteria to enter, and may result in infection in the eye, in the respiratory tract, and elsewhere	It is essential for: Growth Good health at all ages Normal appetite Proper functioning of the di- gestive tract Successful reproduction and lactation Its absence causes: The beriberi, or polyneuritis
	Vitamins	Vitamin	Vitamin B (Anti-neuritic vitamin)

Vitamin

(Anti-scorbutic vitamin)

Good teeth and healthy gums The maintenance of blood Good health at all ages It is essential for:

Fleeting pains in the joints, Insufficient amount may cause: vessel walls

sometimes mistaken for rheumatism

Scurvy

Its absence causes:

Citrus fruits, raw or canned Tomatoes, raw or canned Каж сарьаде Raw peppers Spinach

While they contain only fair

Cod-liver oil, halibut-liver oil, salmon and other fish oils Egg yolk

Ultraviolet rays acting on the skin, either from sunlight or from special lamps (that is, carbon arc, quartz mercury-

Vitamin D is now being intro-duced into some foods which are not naturally good sources (as milk and bread) by irradiation of the food or of some ingredient

fults and vegetables considerably, except in the case of the axed foods such as cirrus fruits and tomatoes. When foods are canned com-Drying and storing foods tend to destroy vitamin C. The canning process tends to re-duce the vitamin-C content of destroyed of the known vita-mins. Exposure to air, long cooking, and the addition of soda in cooking tend toward the destruction of vitamin C Foods canned at home, especially by the open-kettle method, may lose more vitamin C than do commercially Vitamin C is the most readily mercially, air is excluded, and this process reduces the destruction of vitamin C. canned foods

Vitamin D may be somewhat more slowly destroyed by exposture to air than is vitamin A

Ordinary processes of cooking do not easily destroy vitamin D

Since vitamin D is not widely distributed as sources need emphasizing

amounts of vitamin C, raw apples, onions, and turnips, and cooked potatoes may be important sources because they are cheap and plentiful.

It is essential for: Growth

regulating the use of cal-Good bones and teeth (by Good health at all ages the body)

(Anti-rachitic vitamin)

Rickets, which in turn may cause permanent deformities of the bones Its absence causes:

	VITAMIN D	VITAMIN DATA—Continued	•
Vitamins	Functions in the body	Good sources	Effects of various factors on the vitamin
Vitamin G	It is easential for: Growth Good health at all ages Freetion of symptoms similar to those of pellagra, such as digestive disturbance and skin lesions Its absence: Appears to be at least one factor in causing pellagra	Fresh lean meat Liver and kidney Milk, fresh, evaporated, and dried Buttermilk Salmon, fresh and canned Eggs Green leaves Tomatoes Tomatoes Weast germ	Ordinary cooking temperatures and exposure to air have little effect on vitamin G Use of soda in cooking has a destructive action on vitamin G

Egg Preservative British Patent 409,623

Eggs are coated with following: Soft Yellow Paraffin 75 oz. Tallow 5 oz. 20 oz. Boric Acid

Destroying Yeast Spores in Soda Water Bottles

Sonk for five minutes in 1% caustic soda solution at 45° C, and for 10 min-utes in 2% caustic soda solution at

Meat Curing Salt U. S. Patent 1,976,831

Mix together in an aluminum vessel Sodium Nitrite 11/2 lb. Sodium Nitrate Melt while stirring. Pour on metal plate to solidify. Pack in air-tight tins. For treating 100 lb. of beef use 1/4

oz. of above ground into 3 lb. of salt.

English Mustard, Prepared British Patent 412,967

Mustard flour is mixed with cold milk and water with 2% gum arabe and after ½ hour is sterilized by treating at 65-70° C. for 15 minutes, then cooled to 30° C.

Smoked Fish

It is hardly possible to furnish directions for smoking all species of fish, under all the varying weather conditions that will be encountered with the changing seasons. Only the general methods can be given here, as used on a typical variety under average conditions. is intended as a guide, not an infallible recipe. To smoke fish successfully, experiment and use intelligence—altering the method according to the preference of markets (amount of salt and smoke flavor), the variety of fish, and weather conditions.

There are two general methods of smoking fish-hot smoking or "barbecu-

smoking fish—hot smoking or "barbecu-ing," and cold smoking. Any fish may be "hot-smoked" or "barbecued" but the following varieties are some of those to be preferred:

Butterfish Sailfish Kingfish Spanish mackerel Shad Mullet Grouper

Sturgeon is always hot smoked.

Because of the keeping qualities of cold-smoked fish, certain varieties offer market possibilities for quantity production, such as:

Alewife or river herring
Shad
Drum
Mullet
Red snapper
Redfish
Grouper
Kingfish
Robalo or Snook
Squeteague (spotted trout)
Spots

In the first method the fish are laid three or four feet above a fire, and cured at temperatures from 150 to 200° F. The fish are wholly or partially cooked by this method, and therefore, no matter how carefully prepared, or how long smoked, will "keep" for periods of from a few days to a couple of weeks. If fish is to be preserved for any period of time, the cold smoking method should be used. In this process the fish are cured over a low smouldering fire at a temperature of 90° F., or less. The efficiency of the process depends on the drying action of the fire, which must be carried on at a temperature that will not cook the flesh. Fish may be given a short cold smoke, if preservation is intended for a few days only, or cured for several days if it is wished to "keep" them for some time. product is comparable to ham or bacon and should be cooked before using. The same general principles governing smoking, handling, and storing of cured meats should be followed in smoking fish.

A snokchouse for curing small lots of fish may readily be made, following instructions given here. Obtain a box or make one, about 6x3x3 ft. One end, that resting on the ground, should be removed. About 12 in. above this end a false bottom with auger holes at 2-in. intervals is built. This end of the box is set over a pit 2 ft. wide by 18 in. deep.

A trench about 1 ft. wide by 1 ft. deep is dug from this pit for a distance of about 10 ft. The fire pit, a hole 3 ft. wide by 3 ft. long, by 18 in. deep, is dug at the end of this trench, which is then covered by sheets of galvanized iron, forming a chimney for the smoke from the fire pit to the smokehouse. If it is desired to build a more permanent house, terra cotta drain or sewer pipe may be used to connect the fire pot with the smokehouse. Cleats are nailed inside

the box on the sides, the first set about 12-14 in. below the top. The trays for holding the fish, or the ends of the smoke sticks rest on these cleats. A few holes should be bored for ventilation in or near the top of the house.

If mullet or Spanish mackerel are to be smoked, the following process is recommended:

The fish should be split along the back just above the backbone, almost to the tail so that it will lay flat in one piece, leaving the belly portion solid. Clear out all traces of intestines, black skin and blood, taking special care to remove the coagulated blood and kidney just under the backbone. The head may or may not be removed, depending on the individual. If the head is cut off, the hard bony plate just below the gills should be nilowed to remain, as it will be needed to carry the weight when the fish are hung on rods. If it is cut off the fish often pull loose and drop from the sticks.

After splitting and cleaning, the fish should be dropped in a brine made by adding two cups of salt to 4 gal. of water. They are left in this brine 30 minutes to soak out blood diffused through the fiesh. At the end of this time they should be taken out, rinsed, and freed from any remaining traces of blood or other offal. Drain for a few minutes then drop each fish singly in a shallow box of fine salt, "dredging" it about, then picking it up with as much salt as will cling to it, and packing the fish in even layers in a tub or box.

The fish should be left in salt from 1 to 3 hours, depending on weather, size of fish, fatness, and length of time for which preservation is desired. The exact length of time must be determined by the smoker. When the fish are taken out of salt they should be rinsed in brine, scrubbing off all visible particles of salt or dirt. The fish should then be laid on chicken wire drying racks kept out of the direct rays of the sun, but located where a good breeze can reach them. Wire drying racks are desirable as the fish can dry on both sides. One side will remain wet, if laid on boards. The fish should be given about 3 hours drying, until a thin film is formed on the surface, before putting the fish in the smokehouse. If put in immediately after taking out of salt, the fish will be too moist, will require longer smoking will not color and dry as well and will not have as good a surface.

The fish may be placed in the smokehouse on wire mesh trays, or hung on sticks or iron rods. In no case should any two fish touch as this will prevent the drying and penetrative action of the smoke. If hung on rods, more fish may be smoked at one time, and they will smoke better, with a clearer color. Trays, of course, give less trouble. Rods are run through the fish just under the hard boxy plate at the neck, one rod on each side. Thus, each fish hangs from two rods. Twelve or fourteen fish may be hung on a set of two rods 3 ft. long.

The fire should be started an hour or two before the fish are put in the house. It should be low and smouldering. Almost any hardwood or wood other than pine may be used for fuel. Pine or other pitchy woods will give the fish a bitter taste. Some of the woods that may be used in the Southern States, are scrub oak, live oak, hickory, sweet bay, river mangrove, palmetto roots, button wood, and coconut husks. In smoking any one kind of fish, such as mullet, variety of flavor may be obtained through the choice of wood used in smoking. In addition to the woods listed above, orange wood gives a par-ticularly pleasing flavor. Cypress may also be used. The fire should not give off too much smoke during the first 8-12 hours. A dense cloud of smoke should be built up for the balance of the process. The fire must be small and steady. Two short chunks of wood about 2 ft. in length and the thickness of a man's arm are usually sufficient. The fire pit is kept covered with a sheet of metal to drive as much smoke as possible up into the smokehouse, and to keep the fire from burning rapidly. The fire must not be allowed to blaze up. The air should not feel warm on the hand if it is put in the smokehouse. The fish should be smoked for 24 hours, if they are to be kept for a couple of weeks, and for 4 or 5 days if it is wished to keep them for some time. The fire should not be allowed to die out at night or to be built up too large the last thing at night to make it last until morning.

After taking the fish out of the smokehouse dry for an hour or two in the air, then wrap in sheets of waxed paper, sprinkling a little fine table salt on each one, and store in tin or wooden boxes. Keep in a cool, dry place. If signs of mold appear, sponge off with vinegar and give the fish a short smoking for from 3 to 6 hours.

or from a to a manual

Hot Smoking—German Method

The following method is recommended
if it is desired to prepare a hot smoked

fish that can be used immediately without cooking. It will keep without molding or souring longer than other hot smoked fish.

Split, clean, and soak the fish to remove blood, as instructed previously. Then prepare a brine as follows: 2 lb. salt, 1 lb. sugar, ½ oz. saltpeter, 1 oz. crushed whole black peppers, 1 oz. crushed cardamom seeds. Make this up into a 90° brine, that is, one that will float a potato with a 10 d. nail stuck in it. Increase the amount of ingredients according to the quantity of brine you wish to make. The number of spices used can be increased in variety and amount. Various spice mixtures are used.

Put the fish in this brine for a period varying from 2 to 4 hours, depending on the size and thickness of the fish, amount of fat, and the taste of the individual. Some require a less salty taste than others. The exact length of time must be determined by experiment. Rinse off the fish in fresh water, and place on drying racks outside in a cool, shady, breezy place to dry for about 3 hours before putting in the smokehouse.

For the first 8 hours that the fish are in the house, give them a cool smoke in a dense cloud of smoke. Then increase the fire until the temperature is between 130 and 150° F. for 2 or 3 hours, or until the fish have a glossy brown surface. This partially cooks or "hours before storing. Wrap in waxed paper and store in a cool dry place. Do not allow them to come into contact with ice, or store in we toold.

In some cases the fish are brushed over lightly with vegetable oil (usually cottonseed) either just after finishing the cold smoking part of the process, or on taking out to cool. Another method of handling this fish after smoking is to cut the fiesh up into fingers the length of a No. 2 can or pint glass jar. Skin and pack into the can or jar. Then add vegetable oil (cottonseed or olive oil, if you have it) until the spaces between the pieces of fish are filled and there is a layer of oil up to within an eighth of an inch of the top. Sent the cans or jars and store in a cool place such as an ice box until used. Under such conditions it should keep almost indefinitely. As this product is not "sterilized" the cans or jars should be thoroughly scalded before use. In some cases the oil is filled in hot and the containers sealed immediately.

Smoking Fish

Lake Herring and Whitefish

The process of smoking lake herring and whitefish is identical. If the fish are frozen when received at the smokehouse, they are thawed in the open air or better, by immersing and stirring them in a barrel of water of medium temperature. After thawing they are split down the belly to the vent, eviscerated, washed thoroughly, and pickled in butts or barrels, about 4 lb. of fine salt to 100 lb. of fish being scattered among them and sufficient brine of 90° salinity to cover them. Either dry salt or brine alone may be used, the former being preferred in warm weather and the latter during the winter. In case brine alone is used. some dry salt should be placed on top to strengthen the weak pickle floating at the surface. After remaining in the pickle for 10 to 16 hours, according to the strength of the pickle and the flavor desired, the fish are removed and strung on the smoke rods, 10 to 20 fish to each rod, according to its length and the size of the fish.

In stringing, some curers pass the rod through the body immediately below the nape bone, effectively preventing the fish from falling down in smoking, but also marring its apearance somewhat. more usual way is to pass the stick in at the right gill-opening and out at the mouth. Others pass the rod through the head near or through the eyes, and a few pass it immediately back of the throat cartilage. The latter leaves a neat appearance, yet it permits more fish to fall in the smoking process than when the rod is passed through the head or the shoulders. In some houses the smokestick is not passed through the fish, but instead a stiff iron wire, curved in "S" shape, is used to attach the fish to the stick, one end of the wire passing through the fish at the head or beneath the nape bone and the other hung over the smoke-stick. At Grand Haven, and to some extent in Chicago, Milwaukee, and one or two other places, the fish are secured by having stout smoke sticks, about 11/2 in. thick and 21/2 in. wide; in the top of each, and about 1/4 in. from the edge, is driven a row of tacks or small wire nails at intervals of about 3 in., projecting about 1/2 in. above the surface. Ordinary cotton wrapping cord is tied to the wire nail at the end of each stick, and by means of this cord passing around each nail a single herring is held in place between each two nails throughout the length of the stick, the fish being placed with the back of the neck against the stick and the cord passing from one nail around the throat of the fish, entering under the gills on each side, and then around the next nail, and so on to the end. By having the stick of suffiscient width, a row of small nails may be placed on each edge, so as to attach a row of fish at each side. This removes nearly all risk of the fish shifting, and their appearance is not marred by holes through which the smoke-stick has been passed.

Some markets prefer the herring well smoked on the inside and to accomplish this the sides of the abdominal cavity are stretched open by means of small wooden stricks or tooth picks, either one or two sticks to each fish. This permits the smoke to permeate the stomach cavity better and results in a more durable article. In general, the western trade prefers the stomach cavity stretched open, while the eastern markets prefact them without the sticks; but there are exceptions. The smoked lake hearing sold in Washington are mostly extended by means of a small stick, or, in case of large fish, by two small sticks.

The fish attached to the sticks are dipped in fresh water to remove surplus or undissolved salt, loose scales, etc., unless they have been rinsed before stringing, drained, and suspended in their smokelouse 4 to 8 ft. above the floor, and subjected to a gentle smoke for 4 or 5 hours. The door or damper is then closed, the fires spread or built up and the fish cooked for 1 or 2 hours according to the amount of fire, the height of the fish, and the particular cure desired. After cooling, which is accomplished either by opening the doors of the smokehouse or by removing the fish to the outside, they are ready for the trade. One hundred pounds of round fish, or 85 lb. dressed, yield about 65 lb. smoked.

Ordinarily these fish keep one or two weeks, and even longer. Lake Trout and Carp

Smoked lake trout and carp are prepared to a small extent in the manner already described for lake herring or whitefish.

Smoked Fish

Alewives, or River Herring

River herring or alewives are smoked in a number of localities, but principally in Maryland and Virginia.

In preparing these fish in the Chesapeake region they are washed in vats and scaled with a knife as soon as prac-

ticable after removal from the water. They are next immersed over night in strong brine, containing 12 to 14 lb. of Liverpool salt to each 100 lb. of fish, with some dry salt on top to strengthen the weak pickle that rises to the surface. The following morning the round fish are strung on smokesticks, the stick being ntered at the left gill-opening of case of har having or bloaters on the New England and The strings of fish attached to the stick to then dipped in fresh water to rinse the off, and after attached to the stick to then cupped in fresh water to rinse the noff, and after draining; and drying for law hours are suspended in the smokehoot thout 6 or 8 feet above the fire, and an action a dense but cool smoke made of pane and ings or similar material for about the fire afrom becoming too. It is the fire afrom becoming too. It is the cays. Care must be taken to the first from becoming too, in danger and the fish to crack at the care and or possibly to fall from the stick to the core. Prepared in this mannel fleeting the berring will usually keep in good bottom. the true therring will usually keep in-good containing the thesapeake region for 30 days during the spring and to a somewhat less period in the sales As the fish are not eviscerated before smokng the decrease in weight is small, 100 ng the decrease in weight is aman, now be of round falt vielding short 78. Is maked. The wholesale price is about 20 to 22 cents per desen, according to a size and condition.

wo other places the river herring are prepared in the following manner:

The fresh herring are scaled with a knife, gibbed like the pickled herring of Scotland, washed, and pickled for 3 hours in brine, about 20 lb, of Liverpool salt being used for each 100 lb. of fish. On removal from the pickle they are strung on small iron rods, the rod passing through the eye sockets of the fish. drained for an hour or so, and hung in the hogshead smokehouses, in the bottom of which a fire has been made of equal quantities of oak and hickory wood. The fish are dried for a few minutes and then the tops of the hogsheads are covered with old sacks or other suitable material. From time to time the fire is sprinkled with water to produce a vapa and the fish thus exposed to heat, smoke. and steam for about 3 hours, when they are removed and cooled and are then in condition to be eaten. Only oak and hickory should be used as fuel, as other materials do not produce the proper flavor. If the fire becomes too warm it should be smothered with oak or hickory eawdust.

The process of smoking alewives com-

monly employed in the New England States differs from the Chesapeake process in a few minor particulars. The smokers are usually not so careful about removing the scales with a knife, depending generally on the frequent handling of the fish to scale them if cured soon after removal from the water. It is also customary in salting the fish to permit them to make their own pickle, the fish remaining in the pickle for 3 to 5 days. On removal they are soaked in fresh water for 5 to 6 hours and strung on hardwood sticks, the stick entering through the left gill-opening and out at the mouth. They are next rinsed, drained and dried for a short while and suspended in the smokehouse, where they are exposed to a smoldering fire of hardwood and sawdust for 3 to 4 days, when fter cooling, they are ready for sale.

Shad

Shad

The the peake region and at various prints along the coast small quantities of shad are smoked, usually increisely the same manner as already peribed for river herring, or advites. Catfish

Being intended as a substitute, the catfish are smoked in identically the same manner as are sturgeon. The as received at the smokehouse are usually beheaded and eviscerated. They are skinned and cut into small pieces, weighing about 1 or 11/2 lb. each, and are pickled for 6 or 8 hours in tight barrels. This may be accomplished by rubbing the pieces with salt and placing them in the barrel either with dry salt scattered among them, or simply by placing them in the barrel with dry salt or with strong brine. On removal from the brine the pieces are rinsed by dipping in fresh water, to remove slime, surplus salt, etc.; they are then attached to the smokesticks and drained for an hour or so, and placed in the smokehouse where they are smoked for 7 or 8 hours in the same manner as sturgeon are treated. hundred pounds of dressed catfish yield from 65 to 70 lb. smoked, and the product sells usually at about 15 or 16 cents per pound. The total annual product of smoked catfish in the United States probably does not exceed 50,000 lb., and its sale is confined principally to those who are willing to accept a sub-stitute because of its being cheaper.

At several points in the Mississippi Valley the small catfish are smoked whole, like lake herring. They are split to the vent and eviscerated, the head and in some instances the skin being left on.

struck with salt in tight barrels, and smoked for a few hours in the manner described for lake herring.

Eels

Generally the cels are received at the smokehouse fresh, directly from the fisheries, but some are also received frozen from cold storage. In the latter case they are thawed by immersing them in water a few hours or by exposure to the open air. Some smokers "slime" the eels with salt; that is, rub the skin with a small quantity of fine salt to remove the slime therefrom. In dressing, the fish are split from the head to the vent and the viscera removed. It is desirable to continue the splitting down to the end of the tail sufficiently deep to remove the large vein along the backbone, but sometimes this may be pulled out without splitting the fish.

an inch or two beyond the vent. It is smokers, however, give stenden to his item. The cels are immersed in stong brines from 1% to 7½ hours, according to streng of brine, site of fish, and its desired free. This brine should the the fish.

As a York the eels are usually holds for 2 hours, while on the Great Lakes the length of the time is generally about 7 hours. On removal of the fish they are washed, bristle brushes being

they are washed, bristle brushes being used by some smokers, while others simply dip the fish in water for removing the slime and surplus salt. A few smokers throw them in a tub of water and beat them with a net for several minutes to accomplish the same purpose. The eels are next strung on iron or steel rods one-third inch in diameter, the rod passing through the head of each eel, or through the throat cartilage and out the mouth, and hung in the open air for a But if the few hours for drying. atmosphere be moist or the saving of time necessary they may at once be placed in the smokehouse.

In New York, where small brick ovens are used, the fish are subjected to a mild smoke for about 4 or 5 hours until they have acquired the proper color, when the fires are gradually increased and they are hot-smoked or cooked for 30 or 40 minutes. At Buffalo and some of the other Great Lakes ports, the smoking is usually at an even temperature throughout and continues for 6 or 8 hours. Mahogany or cedar sawdust is used in New York for making the smoke, while hickory or white-oak wood is used for

cooking, the latter being preferred. In Washington the cels are suspended in the hogshed smokehouses over a fire made of onk and hickory wood and dued for communities, when the hogshead is covered with sacking and thus hot-smoked for 3 or 4 hours, the fires being sprinkled with water from time to time to produce a hot vapor. The smoking must be capfully attended, for if the baseling the cooking is sufficient to the cooking is sufficient to be superiod or peeled from the cels, when the cell base been split was the result of the cooking is sufficient to be superiod or peeled from the cels, when the cell base been split was the result of the cooking is sufficient to be superiod or peeled from the cels, when the cell base been split was the result of the cooking is sufficient to the cell base been split was the cooking in the cell base been split was the cooking in the cell base been split was the cooking in the cell base been split was the cell and the cell base been split was the cell and the cell base been split was the cell and the cell base been split was the cell and the cell an

been split. The receipt in yeight by dressing and modelly is about 38%, 100 lb, of the been pickled to 78 hours they reliable keep 10 or 12 days; but when they reliable to the been only 2 hours as it issues at New York, they are lithlet to the separation of the receipt of time and the receipt of the receipt

Bels are symen, the before being moved, the process being the same as distributed above, except that let salting and smoking is required, and it is also very difficult to keep them from falling down on the rods in the smoke-

Salting (Including Corning) River

The fish are usually taken from the boats on the day they are caught, but in some cases not until the third or fourth day. All handling of the fish is with scoop nets. When taken from the boats, they are spread upon the wharf for cutting. Sitting on a low inclined seat with his knees on the wharf, the cutter removes the head and belly and scrapes out the roe and viscera, the cut fish being placed in a basket and the roe in a bucket. The fish are then dumped into the washing vats. These are 12 ft. long by 6 ft. wide by 3 ft. deep of 2 in. pine. In some the bottom is inclined about 30° to one side, with a horizontal false bottom of slats above the incline. Sales, dirt and other washings settle dwn in the deep angle of the bottom and are drawn off with the wash water through two flood gates without loss of time. Others still employ flat bottomed vats with resultant loss of time in clean-

ing.

The fish are agitated in the vats (which are kept filled with water) for about 10 minutes to thoroughly wash them and then scooped out with dip nets

into slat cars holding about 1200 fish, in which the fish drain as they are transported to the salting vats. The latter are 10 ft. long by 6 ft. wide and 24 to 30 in. deep built of 2 in. Virginia pine. The salting vats contain saturated brine to a depth of 4 in. As each car of fish is dumped into the brine, additional salt is added, the amount depending upon with which the skilled packer is fully conversant. When full, the vats contain from 12,000 to 15,000 fish (about 4000 lb.). The fish should be roused once each day while striking. After each rousing, the fish are tamped down lightly and top dressed with a thin layer of salt.

Corning

Early in the season most of the packers in the lower Potomac corn their herring for immediate consumption. This method is usually followed for about 6 to 10 days from April 1. The earliest caught, fish are kept in the brine from 12 to 48 hours according to temperature. Fish brine 12 hours when the temperature is from 40 to 50° F. should keep for tep days. After brining, the fish are taken from the vats and spread on the floor, covered with salt and the salt and fish thoroughly mixed, after which they are packed in sugar barrels and immediately shipped to the trade. No fish "are corned after the temperature riscs abous 60° F.

Hard Cure or Tight Pack

Herring intended for storage are keptin the brine for 7 to 10 days according to temperature. At temperatures from 50° to 60° F. 9 to 10 days is sufficient; if from 60° to 70° F., 7 to 9 days will cure them satisfactorily. After the fish are cured, they are taken from the brine and piled on the draining floor to a depth of from 1 to 4 ft. according to available space and allowed to remain there from 4 to 10 days according to the demand for the space. The fish are then weighed or counted (weighing is most accurate) and packed in the barrels, the first layer backs down, the balance backs up with from 2 to 2½ lb. of salt to the layer. A properly packed barrels should contain 160 lb. of fish and 40 fb. of salt

Salted Fish

Considerable trouble has been experienced in salting fish in warm climates. The methods followed commercially in other regions have not produced a

product of good quality, and the directions given generally for salting small quantities, or for the home curing of fish have not always proven satisfactory.

If attempts are made to preserve fish by "pickling" or curing in brine, in a warm climate, the product will either turn "rusty" and sour, spoiling in a short time, or if the quality is good at first the fish soon deteriorates. The best method for curing fish in this region is "dry-salting." That is a combination of salting and drying. If the fish are handled carefully, and directions given below followed closely, a high quality product that will not spoil nearly as rapidly as salted fish now prepared, can be produced. But if instructions are not followed, it is useless to expect much.

In the first place the fish must be absolutely fresh. Do not try to save fish that may be stale, by salting. The fish should be bled, when caught, to drain out all blood possible. Blood decomposes much more easily and quickly than flesh. Fish will keep longer if blood is not diffused through the flesh. They should be thoroughly cleaned as soon as possible. Fish should not be handled roughly in taking out of the net or while in the boat. If fish are piled in heaps, walked on or forked roughly, they will be of inferior quality and spoil much more readily than they would otherwise. Fish should not be left under the direct rays of the sun in an open boat. A tarpaulin should be rigged above the fish.

Mullet and Spanish mackerel are among the best fish for dry-salting, for many reasons, a few of which are: they are split more easily, the loss of weight is less in splitting and cleaning; they are two of the commonest southern fish, and obtained more easily and cheaply. Using this outline as a guide, however, many other varieties of fish, such as grouper, sheepshead, alewives or river herring, spot, croaker, and drum, may be cured successfully, with the resultant product of good quality.

Most fish should be split along the back, just above the backbone, taking care to leave no flesh on it. The fish are split "mackerel style." That is, they must lay flat in a single piece, leaving in the backbone. When the knife is drawn toward the tail it must not go clear through the skin, so that the fish will be in two pieces near the tail. The head may or may not be removed. In splitting Spanish mackerel and other fat fish the backbone is cut out nearly to the tail, where it is broken off. In

cleaning, remove all traces of blood from under the backbone and clear away all the black skin. A wire brush should be used for the blood. "Black skin" is best wiped out by a piece of canvas or gunny sack. If the head is left on, clean out all traces of gills. All cleaning must be done thoroughly and care-

When the mullet or mackerel are cleaned they should be rinsed, then dropped in a tub of light salt brine (2 lb. of salt to 5 gal. of water), the fish should be left here to soak 30 minutes. The principal object of brining is to remove traces of blood from the cut flesh. It also "cuts" slime and is better for washing than water. Never use sea water from around a fish house, dock, or near shore. It is invariably contaminated and increases likelihood of spoilage.

Score with a knife under the backbone and then longitudinally through the flesh on the other side. After the fish have soaked 30 minutes take them out, making sure that each one is properly cleaned. Drain them for 15 minutes. If salted at once the excess moisture will

require more salt.

Use a "dairy fine" ground mined salt. Ordinary sea salt is more apt to cause reddening. Coarse salt is not as good as a fine salt. Pour the salt into a shallow box about 2 ft. square. Dredge each fish in this salt, rolling it about 2 or 3 times and rubbing salt into the slashes. Pick it up with as much salt as will stick to it. Scatter a thin layer of salt on the bottom of the tub or box used for salting. Then lay in the fish in an even layer, flesh side up. Be sure that no two pieces of fish touch without salt between. Scatter a little salt on top. Continue this until all the fish are in salt. Each layer should be laid in at right angles to the preceding layer. The top layer should be weighted down, to keep the fish under the surface of any brine formed. The top layer should also be packed skin side up. Use about 1 part of salt to 3 of fish.

The salting shed should be light, open, airy, and cool as possible. The mullet will have absorbed enough salt for curing purposes in about 36 hours. Mackerel should be in salt about 48 hours. At the end of this time take the fish out of the salt and scrub them in a brine of the same strength as used in cleaning to remove all excess salt and dirt. No traces of salt should be visible on the surface. After draining 15 to 20 min-utes, the fish are ready for the drying racks. These are frames of wood, covered with chicken wire and standing on legs 3 or 4 ft. high.

The drying racks must be placed on dry ground, preferably covered with gravel. Oxidation or rusting sets in immediately if drying is carried on under the direct rays of the sun. But if fish are kept shaded in a breezy location they will dry well with a clear color. For this reason drying is best done in the shade under a roof without walls, so located that as much of a current of air as possible will pass over the fish. The fish are laid out skin side down but are turned 3 or 4 times the first day.

The fish are gathered up and placed under shelter at night to prevent spoilage through dampness. If left spread out in the open at night, they will sour and mold. The time required for drying depends on weather conditions during the drying period, and on the size of the fish being cured. The exact time must be determined by the person curing the fish. For mullet it should average about 4 days; Spanish mackerel, 5 days. The more the fish are dried, the less danger there will be of reddening or rusting. When the surface looks dry and hard, and if the thumb can be pressed into the thick part of the flesh leaving no impression, the flesh can be considered as cured. In weather where air-drying is impossible, or in climates too humid for this process, the following method may are "struck be used. When the fish are "struck through" or have absorbed enough salt for curing purposes, they should be taken

out of salt, scrubbed off in brine, then piled in stacks, flesh side down. These stacks should be heavily weighted down in order to press moisture out of the fish. After 10 to 18 hours in the stack the fish should be repacked in dry salt with the top weighted down, and put in storage in a cool dry place.

Store the fish in wooden boxes lined with waxed paper. Scatter a little dry salt between each layer of fish-about 1 lb. of salt to 10 lb. of fish. Store in as cool and dry a place as possible. If signs of rust or mold appear, scrub the fish off in brine and dry in the air for

aday or two.
Reddening of salted fish is a form of bacterial spoilage caused by the salt used in curing. Contrary to popular belief, salt is not strictly an antiseptic, and certain types of bacteria live and thrive in a salt medium. Salt most apt to be contaminated is that obtained by evaporation of sea water. Several types of salt used extensively in fish curing are apt to be thus contaminated. In salting fish

every effort should be made to use a salt as pure and high in grade as possible. It is advisable to heat salt and bake it thoroughly before using. If, however, reddening appears at any time, all tables and other equipment used in salting should be thoroughly disinfected. Unless every effort is made to keep the salting equipment clean, the use of sterilized salt or other precautions will be useless as the fish can be contaminated through unclean equipment. After curing, the fish should be stored in the coolest place possible, as the salt reddening bacteria grows best at a warm temperature. At first signs of reddening the fish should be removed, washed thoroughly in pure salt brine, and given a few hours careful drying and repacked with a thin layer of dry salt between each layer of fish, using from 10 to 15 lb. of salt to 100 lb. of fish. Reddening is most apt to appear in fish stored in pickle (brine) and held in a warm place. It will remain in good condition longer if packed in dry salt and held in as cool a store room as possible.

Canning Alewives or River Herring; Roe and Buckroe

The following method of canning alewives has proved quite satisfactory. The fish are cut, washed, and placed in the salting vats in the same manner as if intended for salt curing. After 12 to 14 hours they are removed from the vats and washed in an abundance of lukewarm fresh water. During the washing they are trimmed, the balance of the fins and scales being removed. They are then cut to can size and placed in the cans, after which they are processed for 55 minutes at 244° F. for No. 1 cans and 60 minutes for No. 2 cans.

Herring roe intended for canning is collected in buckets as the fish are cut and washed in fresh water in special trays, blood and adhering particles of entrails being removed. The roe is then put in the cans. As it swells considerably in processing, the cans must not be entirely filled. If of the sanitary type, the cans are filled to within about three-fourths of an inch of the top with roe and then filled to the edge with cold salt brine, about 1 lb. of salt to 8 or 10 gal. of water being used to make the brine. The brine is added solely for seasoning. The cans are immediately capped and placed in the processing baskets. If solder-top cans are used, the filled cans are placed in the exhaust box. Upon removal from the exhaust, the necessary

air space is provided for by pressing the roe down with a plunger. Material clinging to the groove where the solder is to be applied is removed with a brush and the cans are capped and tipped. The canned roe is processed in a closed kettle for 45 to 55 minutes at a temperature of 240°-245° F. The milt roe may be canned in the same manner as the roe except that the cans can be more completely filled, as this product does not swell in the processing. As the quantity of brine used in this case will be somewhat less, it should be made correspondingly stronger.

Note: In canning the fish, they should be drained of superfluous water before they are placed in the cans, and no water added to can contents. That the fish may retain their shape in the can and stand transportation, the cans should be well filled. The shrinkage of the fish in processing must be taken account of in filling the cans.

Canning Clams (Alaska)

The first operation is the removal of the clams from the shells. This is done by immersing them in boiling water, either in vats especially designed to receive the wire baskets in which the clams are placed or the clams are passed through the water on an endless belt. After remaining in the water several minutes they are thrown on a table and the shells fall away from the meat. The clams are then passed on to women workers, who open the stomachs and necks, remove the sand and sediment therefrom and sever the black part of the neck. The cleansing process is continued by placing the meat in a cylindrical perforated washing machine, which revolves automatically half a turn both ways in a tank filled with water. Any sediment that may have remained after the hand operations were completed is thus removed. The clams are now ready to be canned and are taken directly to the filling tables if whole clams are packed, or to the grinder if the minced variety is desired. The cans are filled by hand with both meat and juice, after which they pass through the topping and sealing machines and are sealed. The process is completed by cooking the canned product in retorts at a tempera-ture of about 245° F. from 1 to 11/2 hours, depending upon the size of the container used. The juice which is thrown off in the process is used in pre-paring the finished product, the surplus being sealed in cans.

Anchovy Paste

Anchovy paste from sprats may be made as follows: Sufficient for a peck of sprats—2 lb. common salt, 3 oz. bay salt, 1 lb. saltpeter, 2 oz. prunella, and a few grains of cochineal, pounded well together in a mortar; into a stone jar place first a layer of fish, then of the pounded ingredients, and so on until the jar is filled; press them hard down and cover closely. After 6 months they will be ready for use.

Note: Persons using such preservatives as saltpeter should consult the Bureau of Chemistry, Washington, D. C., to determine whether they are using an amount in excess of that held to be proper under existing law.

Anchovy Butter

Take 1 part of anchovies which have been beaten to a paste, and pass through a sieve; add 2 parts of butter, and spice to suit. Cayenne pepper or paprika may be used to advantage.

Anchovy Essence

Anchovy essence can be made with their canned or bottled anchovies. Take the fish, and rub to a pulp in a mortar, and then pass through a fine sieve. To ¼ lb. of anchovies add ¼ lb. of water; boil for 15 minutes, and strain; then add ½ oz. of salt and ½ oz. of flour, and the pulped anchovies. The mixture is allowed to simmer over the fire for 3 or 4 minutes. After the preparation is cool add 2 oz. of strong vinegar. The product should be bottled, in small bottles and tightly corked and covered with bottle wax.

Anchovy Paste

Prepared by taking 1 lb. of anchovies, 1 lb. of water, and 2½ oz. of salt and 2½ oz. of four; add a small quantity of cayenne pepper (say ½ oz.), a small quantity of grated lemon peel, and ½ oz. of mushroom catsup.

Anchovy Sauce

Take 3 or 4 anchovies, and chop them fine; add 3 oz. of butter, 2 oz. of water, 1 oz. of fungar and 1 oz. of flour. Melt the butter over a water bath, add the water and the vinegar, and lastly the flour and the anchovies; stir until the mixture is thick, then rub through a wire sieve. This preparation should be kept on ice, and will not keep indefinitely.

Mushroom Catsup

Upon a suitable quantity of the fresh mushrooms sprinkle salt (about 1 to 4 of the fungi), and after 3 days squeeze out the juice. To every gallon of juice add black pepper, ginger and cloves, of each ½ oz.; pumento, 2 oz.; mustard seed, 2 oz.; and a sufficient quantity of salt. Boil for 5 minutes and set aside to settle. Strain after 7 days.

Christiana Anchovies

In the preparation of Christiana anchovies many methods and flavoring ingredients are used, depending on the skill and ideas of the curer and the markets for which the preparation is intended. The following is one of the most popular processes:

The fresh sprat or anchovies are immersed in brine for 12 or 18 hours, 15 lb. of Liverpool salt being used for each 100 lb. of fish. On removal, the fish are drained in a sieve and then loosely packed in a barrel, with the following ingredients, which have previously been finely crushed and well mixed: 4 lb. of Luneburg salt, 6 units of pepper, 6 units of sugar, 6 units of English spices, 1 unit of cloves, 1 unit of nutmeg, and 1 unit of Spanish pepper. The anchovies re-main saturated with these ingredients for 2 weeks, when they are repacked tightly in kegs or barrels, being carefully arranged in layers, with the backs downward. A quantity of the ingredients above mentioned is sprinkled over each layer, with the addition of a few cut bay leaves or cherry leaves. At the bottom and the top of the package is placed two whole bay leaves, but before the top leaves are laid on, brine is poured over the fish. The barrels or kegs are then coopered and rotated daily for the first few days, and after that every other day for 2 or 3 weeks.

The following process is also used to some extent.

The fish are salted for 24 hours and next immersed in sweetened water, 20 parts of water to 1 part of sugar being used. The fish are then packed with a mixture of Luneburg salt with 90 units or parts of allspice, 60 units of pulverized sugar, 19 units of whole peppers, 15 units of cloves, an equal quantity of nutmeg or mace and of hops (Origanum creticum), and some bay leaves.

The following is a choice method of preparing "Matjeshering" in Germany:

Fresh full herring, both spawners and milters, are well washed, and the gills.

stomach, and intestines are removed in such a way as not to necessitate cutting the throat or abdomen, this being accomplished by pulling them through the gill flap. The fish are next immersed for 12 or 18 hours in a 7% solution of whitewine vinegar, from which they must be removed before the skin becomes flabby and be wiped dry and covered with a preparation composed of 2 lb. of salt, 1 lb. of powdered sugar, this quantity being sufficient for 75 herring. The fish are then packed in a barrel which is sealed. When there is not sufficient brine to fill the barrel, additional should be made of 1 part of the above mixture and 4 parts of water which has been boiled.

Spiced herring (Gewurzhering) are prepared in Germany in the manner above described, with the addition of spices mixed with the salt. The spices commonly used consist of 1 part of Span-ish pepper, 5 parts of white pepper, 4 parts of cloves, 21/2 parts of ginger, an equal quantity of mustard, and a particle of mace and of Spanish marjoram, with a few bay leaves scattered between the

Smoked Pork Sausage

Formula .- Meats: 100 lb. strictly fresh pork trimmings, 85% lean and 15% fat.

Seasoning:

21/2 lb.
10 oz.
4 oz.
1 oz.
⅓ oz.
2 oz.

Nutmeg and ginger may be omitted and sage substituted. Some classes of trade prefer this product with only salt, pepper, sugar and nitrate of soda in the

seasoning formula.

Processing.-Inspect pork trimmings to see that they are fresh and lean. It may be necessary to re-trim, removing blood clots, gristle and hair. Proportion of fat and lean should be closely watched since fat has a tendency to render out in the smokehouse and soften the product. Grind pork through 5/32 or 4 in plate of the hasher, first making sure knives and plates are sharp. Some packers use a rocker entirely for pork sausage.

Place meat in mixer and add seasonings. Mix seasonings and meat for about 5 minutes or until ingredients are thoroughly intermingled. At the time

crushed ice (not more than 7 or 8 lb. per 100 lb. of meat) may be used.

Stuffing .- After seasonings, meat and ice are thoroughly mixed, the product goes to the stuffing bench where it is stuffed in medium hog casings. Link in double links, 31/2 in. in length, knotting ends of casing to prevent meat dropping on truck or floor. Trim off all scrap ends of casings on the outside of knot, but be sure scraps do not get mixed in with the meat.

Carefully puncture casings to prevent air pockets between casings and meat. Sausage must be hung on a truck as fast as it is linked. When truck is filled, put it under an overhead cold water spray for several minutes to thoroughly remove grease and sediment from outside of casings.

Scrap meat on the bench should be handled promptly and mixed with meat stock in the truck. It should not remain on bench for any length of time as it

deteriorates rapidly.

Cooling.—After stuffed sausage has been sprayed it is taken to cooler and spread on trucks or in hanging sections and allowed to hang overnight at a temperature of 36 to 40° F. Product is removed from cooler the next morning and allowed to remain in natural temperatures for about 2 hours.

Smoking.—Then it is placed in the smokehouse at a temperature of 115 to 120° F. and carried at this temperature for about 3 or 4 hours. It does not re-

quire a heavy smoked color.

After smoking it is placed in the cooler at a temperature of 45 to 50° and allowed to hang for 2 to 3 hours until thoroughly cooled. Then it is packed in cartons if it is to be shipped promptly. This product should be manufactured only as needed.

Pork Sausage

Meats:	
Cali Butts	45 lb.
Selected Ham Fat	55 lb.
Seasoning:	
Salt	1¾ lb.
Fine White Pepper	7 oz.
Fine Sage	2% oz.
Cardamom	1/4 oz.
Savory	⅓ oz. ¾ oz.
Marjoram	1/8 OZ.
Ginger	1 oz.
Sugar	3 07

Put ham fat on rocker with 3% ice for 8 minutes, then add seasoning and seasoning is added a small quantity of lean meat and rock for 10 minutes more, making 18 minutes altogether. Meats are all fresh and in small pieces. When rocking is finished fat must have the appearance of half the size of a coffee bean.

Another meat formula for breakfast sausage is as follows:

Shoulder Fat Pork
Trimmings 25 lb.
Pork Butts Trimmed 25 lb.
Lean Pork Trimmings, 40%
Lean (No Belly Trimmings) 50 lb.

"Skinless" Pork Sausage

Sausage meat for this product is stuffed in "NoJax" or similar casings, linked usually in about 4½-in. lengths, and handled and peeled in same manner as skinless frankfurts.

Following are two formulas for "skinless" smoked sausage: For formula No. 1 use, per 100-lb.

batch:
Lean Pork Trimmings, Cured 60 lb.
Regular Pork Trimmings,

Cured 20 lb.
Lean Beef, Cured 20 lb.

Pork is ground through 1/2. in. plate. Chop beef very slightly so it will act as binder and then add to pork in mixer. Care should be taken that no excess moisture is added as it will produce sourness in finished product. Mix well and season with proper amounts of saft, pepper and whatever other seasonings are desired.

Ready prepared seasonings or specially prepared seasonings as manufactured by reputable firms will assure convenience and uniformity in making this product.

Stuff mixture in 1½-in. "NoJax" or similar casing. Smoke in a cool house for 3 hours at 130° F. Then cook at 160° F. for about 10 minutes. Cooking is usually done in a steam house to prevent smearing. Sausage should be placed before a fan following cooking to dry off casing. This aids in prevention of any mould or bacterial growth.

Formula No. 2 uses, per 100-lb. batch:
Cured Pork Cheeks 50 lb.
Cured Regular Pork Trimmings 50 lb.

This formula is prepared in same manner as No. 1. Product must not be chopped too fine or cooked too much to prevent pork from becoming smeary and spoiling its appearance. Sausage should not be peeled or packed in boxes until ready for shipment.

Italian "Hot" Sausage

A good formula for this product is as follows:

Beef, Free of Sinews 60 lb.
Pork Trimmings (Half Regular and Half Lean) 40 lb.

Chop meats through the 1-in, plate and mix with following:

No. 3 Can Pimientos, Juice and All, Chopped to a

Paste 1
Straight Ground Chili

Pepper 1½ lb. High Grade Paprika 1 lb.

If fresh meat is used in making the product 2 lb. of salt should be added. If meat is cured, the additional salt is not necessary. Also add:

Ground Caraway 1 oz.

Ground Caraway 1 oz.
Coriander 2 oz.
Celery 1 oz.
Nutneg 2 oz.

After a thorough mixing, run the product through 342, 1/6 or 3/6-in. plate, depending upon fineness or coarseness of meat desired.

Stuff mixture in hog or manufactured casings, linked 6 to pound. This allows serving two sausages on average plate lunch. Put sausages into cook tank with water at 160° F. and let temperature drop back to 150°. Cook for 30 minutes or until an inside temperature of at least 137° is obtained.

This sausage can be smoked right after it is stuffed, smoking for half an hour in a cold smoke.

Any good bologna or frankfurt meat formula can be used for this sausage, cutting the meat coarser if desired and seasoning highly, with seasonings such as those suggested in the above formula.

as those suggested in the above formula.

Another meat formula which might be used is as follows:

Beef Chucks 70 lb.
Pork Cheek Meat 20 lb.
Back Fat Trimmings or
Shoulder Fat 10 lb.

Grind beef and pork checks through the %-in. plate; back fat trimmings through %-in. plate.

Head Cheese

The following formula can be used to make an attractive product which is strictly a head cheese.

Meata:

S. P. Pork Tongues	60 lb.
S. P. Pork Snouts	20 lb.
Pickled Pork Ears	10 lb. "
Pickled Pork Rinds	10 lb.

Seasoning:

Ground White Pepper 4 oz. Caraway Seed 2 oz. Marjoram 14 oz. Ground Cloves 14 oz.

Prepared seasonings may be used if desired, such as those made by reputable seasoning manufacturers, to facilitate convenience in handling and uniformity of product.

of product.

Cook each kind of meat separately in nets, at 212° F. as follows:

 Snouts
 1½ hr.

 Rinds
 2 hr.

 Ears
 1½ hr.

 Tongues
 1¾ hr.

Grind skins through 1/8-in. plate of hasher. Snouts and ears should be put through 1-in. plate. These should be rinsed several times with warm water to remove surplus sediment and fat.

Remove gullet bones from pork tongues after cooking. Cut each tongue crosswise 3 times, making 4 approximately equal pieces, so that tongues will pass through valve of stuffing machine.

Put all meats togother in a box truck, adding seasoning, jelly water and salt to taste. Not much salt will be required, as all meats used are pickle-cured. Use the hot meat liquid in which meats were cooked, and mix thoroughly.

Stuff tight in hog stomachs or manufactured casings. Fasten carefully and cook 1½ hours at 170° F. Wash cleun and put into cooler at about 36°, or keep in ice water, to chill thoroughly before packing. Product must be clean and free of grease before packing and sale.

Some sausage makers add pimentoes or green peppers to give eye and taste appeal to their head cheese.

Curing and Smoking Frankfurters

Curing is best done by dry-curing hashed ments, by emulsion curing, or by a combination of both. In dry-curing hashed trimmings use per 100 lb. of meat, 3 to 3½ lb. of salt. Nitrate or saltpeter should never exceed 3 or., while nitrite should never exceed 4 or. per 100 lb. of meat. A mixture of these is still better, namely ½ to ½ or. of nitrite and 3 to 2½ or. of nitrite and 2 to 2½ or. of nitrate or saltpeter. The same proportions hold for the emulsion cure. Dry cured hashed trimmings may be used after 2 to 3 days, but they may also be kept 7 days. Emulsion cured meats are put through the fine cutter, and so cure rapidly. Thus they must be used promptly.

Every sausage maker knows that good

muscle meats make good sausage and that cheeks and other such meats do not make sausage of quite as high a class. Less ice should be used in the summer than in the winter. For winter about 60 lb. of ice can be used per 100-lb. block of meat, but only 40-48 lb. should be used in the summer for first grade frankfurters. Less ice can be used with second and third grade frankfurters.

Frankfurters should be properly cured before smoking. If the emulsion cure is used in whole or in part, the meat or the sausage should be held a while for the cure to develop. Part of this may be done in the smokehouse. The smoke should start cool (about 90° F.) and finish at 130-135° F. for Vienna style frankfurters. For other smoked sausage the finish may be at up to 175° F. Cooking should follow promptly and the two operations should really be considered as one. Vat water should be 160°-165° F. while in the spray cooking process the water may be 180° F. Cooking should proceed until the temperature at the center of the meat is at least 140° F. while 148° F. gives better color and many believe it gives better texture and flavor.

German Ham

Since these hams are not cooked before they are eaten, all packers operating under federal inspection must follow B.A.I. rules for uncooked pork in making them. The way they make them in Germany is as follows:

Only hams with a pink meat color are chosen. They should weigh about 18 lb., and are long cut with some of loin end on. Hip bone should be removed.

For curing use a mixture of 25 lb. of salt and 4 oz. of sodium nitrate, or prepared curing mixture. This mixture is rubbed into the ham, especially the skin side, for about 5 minutes. Press some of salt into leg bone at cut. Place hams in a vat, and on each layer add enough of curing mixture so that all parts are lightly covered with it.

When vat is full it should be covered with boards with a weight on top Curing will take 28 days at not less than 38° F. Repack 3 times during this period, so that top layer goes on bottom. Rub hams over again at each repacking.

At end of 28 days take hams out of vat and lay on floor in same temperature for 14 days, sprinkling curing mixture very lightly between each layer. At end of this period wash hams in warm water and hang in dry-room for 2 to 3 days.

Then smoke in a very cold smokehouse for not less than 6 weeks. In Germany these hams are sometimes smoked for 6 months.

Careful handling in cure will yield a tender product. Packers preparing this type of ham for the first time should cure only a small batch. In this way they can watch smoking and curing closely.

Bologna

To make and cure bologna in the silent cutter one sausage expert advises the use

i an iresi meats, as ionows:		
Beef Chucks	70	lb.
Pork Check Meat	20	lb.
Pork Back Fat Trimmings		
or Shoulder Fat	10	lb.

Grind beef and pork cheeks through the 1/4-in. plate; back fat trimmings through 3/8-in. plate. Put beef and pork cheeks in silent cutter and add cure, as follows:

Salt	3 lb.
Sodium Nitrate	2 oz.
Nitrite of Soda	1/4 oz.
Sugar	6 oz.

· and proceed as if using cured meats.

Add ice and water up to 20 lb. per 100 lb. of meat, and chop for 3 minutes. Then add pork back fat and seasonings:

Ground White Pepper	6 oz.
Ground Allspice	1 oz.
Coriander	2 oz.
Ground Nutmeg	2 oz.

Chop 2 minutes more. Then put in a meat truck or pans not over 6 in. deep, and hold in cooler at 36 to 38° F. over night or about 12 hours. Next morning stuff and let sausage hang in room temperature for 1 to 2 hours. Then smoke, slowly at first, gradually increasing temperature from 120 to 145° F. Cook 45 minutes at 160° F.

This method has the advantage of saving a lot of labor, decreases inventory holding and produces a fine, tacky product.

Non-Discoloring Salami

Discoloration is usually due to curing methods. To make either hard or soft salami, meat should be cured as follows:

Use 2% oz. of sodium nitrate for each 100 lb. of meat. Beef requires 3 lb. of salt and pork 21/2 lb. for each 100 lb. of meat cured. Run meat through 1-in. plate with above curing materials and then cure for at least 8 days at a tem-perature of about 40° F. Then place in mixer, add 9 oz. sugar and 6 oz. of pepper, and mix pork and beef together. Grind mixture through desired plate, either 14-in. or 36-in.

Stuff material tightly in large hog bungs, beef middlings or manufactured casings, as tightly as easing will stand. Hang in a dry chill room for 4 days. Then remove to sausage kitchen and hang for at least 6 hours so it will be raised throughout to room temperature before it goes to smokehouse. It may either be smoked through or smoked 12 hours and finished in cooker.

"Smoked through" means about 24 hours at slow smoke at 90 to 100° F. Then gradually raise temperature to about 140° so that product will have a 137° temperature at center when finished. Remove from smokehouse and rinse off with cold water; allow it to cool before

placing in chill room.

Meat from full grown animals should always be used for hard sausages, such as jumbo shoulder trimmings and large beef chucks with all sinews removed.

A good meat formula for salami is as follows:

Lean Pork Trimmings	50 lb.
Medium Lean Beef Chucks (Free of Sinews)	35 lb.
Back Fat	15 lb.
These ments should be cured	necordin

to directions given previously.

The product may be sessoned with:

Im product may be	pourout	with.
Crushed Garlic	11/4	oz.
Sugar	9	oz.
Brandy Flavoring	5	oz.
Ground Anise Seed	1	OZ,
Ground Cardamom	1,4,	oz.
Maple Flavor	3 -	than.

Coloring and Flavoring for Meats British Patent 425,567

Hæmoglobin, Defibrinated 100 oz. Sodium Nitrite oz. Sodium Nitrate 13/3 oz. Water 100 OZ.

Stir well for a few hours. Spray dry or vacuum dry. 1% of this product is used on meats.

Preserving Color of Meat U. S. Patent 2,009,587

By coating freshly cut meat surfaces with a glycerin-gelatin-water solution containing a small amount of essential oil, the natural fresh color and appear ance of the meat is maintained.

Various essential oils, such as oil of cloves, may be used, or a mixture of oil of black pepper, coriander and allspice.

One typical formula for such a solution that has been found satisfactory consists of 57% water, 25% glycerin, 19% gelatin, and substantially 0.1% of essential oil. This solution may be applied with a brush or spraying device on cloth placed over the cut surface of the meat.

The entire piece of meat may be wrapped in fabric such as export beef cloth or the fabric may be applied only on the cut surfaces. The coating is then allowed to congeal. The glycerin, being hygroscopic, preserves the gelatin in a flexible condition, thus avoiding cracking. The essential oil acts as a germicide, while the gelatin acts as a hermetic seal.

Export beef cloth has been found superior to other fabrics for keeping the preservative solution in contact with the meat.

Preserving Vegetables and Fish Dutch Patent 34.553

A procedure for keeping fruit, vegetables, etc., in a fresh condition has been devised. It is especially adapted for the prevention of mold, fungi, and other micro-organisms developed during storage. The procedure consists in rendering the air of the storeroom slightly alkaline, so that moist indicator paper showing a color change at pH = 7.5 is affected on introduction into the chamber. In order to render the storeroom alkaline, materials which furnish volatile, alkaline substances are burnt slowly.

Preventing Mold on Stored Meats

The humidity of the cooler should be 90 to 92% and the temperature 38-39 * F. Ozone is introduced until it is present in 2.3 to 2.7 parts per million. This is continued for 2 hours and again for 2 hours after a lapse of 12 hours. After an interval of 30 minutes, workmen can safely enter the room,

INKS AND MARKING COMPOUNDS

Ink for Document	8	Pour this into:		
Gallic Acid	5 φ.	Water	180 g.	
Borax	5 g. 0.5 g.	Indigo Carmine, Paste in	200 g.	
Pierie Acid		Water	36 g.	
Ammonia		Wood Vinegar, Crude	15 g.	
	20 g.	Dye for Black Writing: per		
Water	50 g.	Ink add:	1000 66.	
Dissolve with warming and	stirring.	Phenol Blue 3F	10	
Water	50 g.	Ponceau RR	1.8 g.	
Caustic Potash	1 g.		1.2 g.	
Boil and stir the mixture	- 6	Amhne Green D	1.2 g.	
brown, let stand warm for ar	tinth pare	No. 2		
add the following dissolved by	nour, then	Indulible Luk Stable Against V	Indelible Ink, Stable Against Water, Oil,	
3	•	Alcohol, Alkalı, Oxalic Acid,	Chloridan	
Water	200 g.	a. Shellae	4 g.	
Borax	1.5 g.	Borax	2 g.	
Shellac	3 g.	Water	30 g.	
Anıline Blue	4 g.	Boil till dissolved,	30 g.	
-				
Non-Corrosive Writing	Ink	b. Gum Arabic previously	≅ g.	
Gall Nuts	28 g.	Water dissolved	4 g.	
Aniline Blue	6 g.	Mrx a and b, boil, filter, cool	l. ndd	
Ferrous Chloride	30 g.	c. Indigo Carmine to desire		
Glycerin	2 g.	Note: Just traces of sul		
Hydrochloric Acid	30 cc.	hydrochloric acids or salt mak	muric or	
Arsenic Acid		debble.	e ink in-	
Phenol	1 g.	denoie.		
Water	1 g. 1000 l.			
		Ink for Writing on Cellu		
Powdered Writing In	ıks	Ferric Chlorido	10 g.	
Formula No. 1		Tannic Acid	15 g.	
Gallic Acid	10 ~	Acetone	100 cc.	
Ferric Sulphate	10 g. 10.7 g.	Dissolve the ferric chloride	in a por-	
Oxalic Acid		tion of the acetone and the ta		
Soluble Blue Dye	2 g. 3.5 g.	in the remainder and mix the		
•	J.J g.	any pen.		
Formula No. 2		1		
Gallic Acid	10 g.	50 1 7 7 7 7 1		
Ferrous Sulphate Crystals	15 g.	Black India Ink		
Tartaric Acid	1 g.	a. Bornx	0.3 g.	
Soluble Blue Dye	3.5 g.	Shellac, Wax-Free	1.5 g.	
-		Water (Boiling Hot)	4 g.	
Indelible Inks		b. Black Tar Dye, Water-	_	
Formula No. 1		Soluble	0.1 g.	
		Water	4.1 g.	
a. Chinese Gall Nuts,	==0	Mix cold.	-	
Powdered	750 g.			
Water, Hot	3000 g.	Non-Congulating India 1	n.b	
Stir, keep standing 2 days,	then press	" "		
out extract; add to the extract		Japanese Patent 110,28	52	
b. Ferric Sulphate in Water	г, і	Glue (Previously Heated at		
(sp. gr. 1.48)	48 g.	120° C. for 3 hr.)	30 oz.	
Solution, Saturated, of		Urea	10 oz.	
Oxalic Acid	18 g.	Potassium Nitrate	60 oz.	
	18	39		
	-			

Urotropine Carbon Black	10 oz. 60 oz.	Ink for Writing on Car	•
Water	1000 oz.	U. S. Patent 1,98	0,123
This ink will not coagula	te at tempera.	Titanium Dioxide	1 oz.
tures down to -30° C.	oc at tempera	Mineral Oil	2 oz.
		Mineral Spirits (Naphtha	a) 4 oz.
Silver Glow In	nk 1 oz.	Carbon Paper I	nk
Mercury	2 oz.	French Patent 774	
Grind together until liqu	id: then grind	Cottonseed Oil	, 1 lb.
with 1 pint of 2% gum a	rabic solution.	Prussian Blue	1 lb.
When used as an ink the	writing will	Carnauba Wax	2 lb.
resemble silver.		Paraffin Wax	2 lb.
		Ozokerite	1/2 lb.
Marking Ink for Chemic	al Porcelain	Octadecyl Alcohol	1 lb.
Cobalt Oxide, Black Cor	n-		
mercial	18.8 g.	Transfer Ink	
Bismuth Subnitrate	1.2 g.	U. S. Patent 1,990	193
Grind these together thou	roughly with	· ·	•
Turpentine	15 cc.	Carnauba Wax	3 lb. 2 lb.
Dresden Thick Oil	15 drops	Boiled Linseed Oil Caustic Soda	0.375 lb.
Mark the porcelain with	a pen, heat	Pigment	to suit
slowly to evaporate the liq	uids, and then	1 1gmcm	to bare
ignite strongly. The porce	lain apparatus	m It Detail	T.1.
is then ready for use.		Thermographic Printi	
	•	U. S. Patent 1,992	2,016
Ink Erasing Fl	uid	Paracumarone Resin	100 lb.
An alkali hypochlorite, f	irst applied to	Dibutyl Phthalate	50 lb.
the ink to be removed; f	ollowed by an	Butyl Stearate	50 lb.
application of dilute acid	, will remove	Drier	21/2 lb.
iuk from paper.			
T 1 4 01 D 11		Rotogravure In	
Ink for Glass or Polis		French Patent 776	5,825
Sodium Silicate	2 oz.	Ethyl Cellulose	5 lb.
Liquid India Ink	10 oz.	Alcohol	155 lb.
Use on clean surface wi	thasteelpen.	Alcohol Soluble Dye	40 lb.
Ink for Glas	8	Offset Printing I	nle
Turpentino	20 g.	U. S. Patent 1,989	
Venice Turpentine	6 g.		,
Shellac	10 g.	Pigment	34.4 lb.
Mastic	2 g.	Linseed Oil	21.5 lb. 33,2 lb.
Lampblack	6 g.	Varnish Castor Oil	33.2 lb. 2.2 lb.
The lampblack is added		Stearin	3.7 lb.
the mixture of other ingr		Turpentino	5 lb.
ink is very efficient for wi photographic plates and la			
photographic places and la	meen since.	T. A N D. ! . 4 !	T 1.
a. 11 135 11		Intaglio Printing	
Stencil and Markin	~	U. S. Patent 1,962	•
U. S. Patent 2,00	12,939	A pigment is used with	the following
Shellac Solution (4 lb. p	er	binder:	0 11
gal.)	32 oz.	Rosin Countie Potech (100%)	2 lb. 1.6 lb.
	.3–6 oz. .0–2.3 oz.	Caustic Potash (10%) Casein	0.1 lb.
Lampblack or Chrome	.0-2.13 02.	Ammonia (sp. gr. 0.91)	0.1 lb. 0.24 lb.
Yellow 5	.7-8 oz.	Turpentine	0.2 lb.
	-167 fl. oz.	Water	4 lb.

INKS AND	MARI	ING COMPOUNDS	191
Lithographic Bronze Printing I	nk	Newspaper Ink	
Varnish		Pit Coal Tar (0.85-0.89	
German Patent 604,019		Density)	1 kg.
Polymerized China Wood Oil 10	lb.	Linseed Oil Boiled with Lithurge	4 kg.
	lb.	or	-
Carnauba Wax 1	lb. lb.	Linseed Oil-Colophony Varn	ish 4 kg.
Polymerize China wood oil at 240- C., add linseed oil and heat to 20		Pyroxylin Printing 1	'nk
for 2 to 3 hours. Cool and add	1 the	Ethyl Oxalate	10 lb.
carnauba wax and turpentine. About 9 lb. of above is stirred wi	41. 1 <i>a</i>	Nitrocellulose (1/2 sec.)	3 lb.
to 18 parts bronze powder.	tu 10	Dye (Basic)	2 lb.
		or Pigment	2 lb.
Printing Lacquer		- I I I I I I I I I I I I I I I I I I I	2 ID.
U. S. Patent 1,996,846		Typographic Ink	
Nitrocellulose about 10 parts,	ester	Red Yacea Gum, Powder	15 g.
gum about 25 parts, xyloi abou parts, fenchone about 30 parts, di phthalate about 5 parts and pig	t 30	Borax Solution, Boiling	4 g.
parts, renemone about 50 parts, or	ment	Glycerin	1 g.
about 25 parts relative to the total of	fthe	Gum Arabic	2 σ.
other ingredients.		Soluble Nigrosine Water	5 g. 73 g.
State of the state		,	10 g.
Solid Color for Rubber Printing B	locks	Water-Soluble Printing	Tule
Hansa Yellow 200			
Alcoholic Shellac (50%) 50	g.	Glycerin Gum Arabic	100 oz. 50 oz.
Borax 50 Water 250	g.	Water Soluble Dye	10 oz.
	-		
Ink for Rotary Press		7141 11 - 0.1 - T	
Pit Coal Tar (Density		Lithographic Color I	
0.85-0.89) 100	g.	Glycerin	10 g.
Treat warm with:		Coparba Balsam Venice Turpentine and	20 g.
	g.	Sandal Wood Oil	5 g.
then neutralize with stirring by	Boda orida	Petroleum Oil	2.5 g.
Ash. Deodorize with calcium chl and hydrochloric acid.	orace	Pine Turpentine	2.5 g.
Above Tar plus		Alcohol Manganese Dioxide	5 g. 2.5 g.
	g.	This mixture, prepared on	
•	g.	bath, is thinned with	VIIO #440.1-
To this liquefied and cleared va	rnish	Chloroform	16 g.
	g.	Ether Ammonia (28° B6.)	16 g.
to obtain: Black, brown or violet coloration		Ammonia (20 De.)	31 g.
Alum		Lithographic Ink for Repre	oductions
Copper Sulphate		Resin, Damar	12 g.
Potassium Bichromate	1	Petroleum Oil	2.8 g.
Finally mix with		Glycerin	32 g.
Lamp Black 10	g.	Linseed Oil Varnish Color	24 g. 2-8 g.
Typographic Ink for Newspaper	.		2-8 g.
		Fusible Lithographic	In k
Colophony Tar 37 Rosin Oil, Rectified 40	g.	Damar	50 oz.
Thinner: Petroleum 20	g.	Kerosene	100 oz.
Filter hot.		Pigment	100 oz.
	•		

Fine Lithographic Ink	Medium Varnish (for Inks)
Asphalt (Gilsonite plus 60%	Rosin Oil 95 g.
of Rosin Oil plus 70 to	Crude Linseed Oil 35 g.
120% of Rectified Tar) 15 g.	Sulphonated Rosin Soap 7 g.
Pit Coal Tar 30 g.	Colophony 40 g.
Paris Blue 2 c.	20 6.
Bone Black 3 g.	Evanescent (Invisible) Inks
Lamp Black 23 g.	Formula No. 1
To get a typographic ink, increase the	
proportion of tar, and reduce sensibly	Cobalt Chloride 1 dr.
the proportion of the color.	Mucilage of Acacia 1 dr.
the proportion of the color.	Distilled Water 1 oz.
	Dissolve. The writing becomes blue
Typographic Ink for Prints	when the paper is heated, and disappears
	again on cooling.
Colophony Tar 95 g. Rosin Oil (Medium, Neutral,	No. 2
Rectified) 50 g.	*Oxalomolybdic Acid 15 gr.
Linseed Oil, Light 13 g.	Distilled Water 1 oz.
Elimeted Oil, Eligite 10 g.	
	Dissolve. Write with this in a dull
Lithographic Inks with Oil-Varnishes	light. When exposed to sunshine, the
Thickened by a Resin	writing appears blue; when wetted, the
	blue changes to black.
	* Made by dissolving Molybdic Acid to saturation in a hot solution of oxalic acid, and collecting the crystals on cooling.
Varnish, Medium 40 g. Soda Ash 2.8 g.	collecting the crystals on cooling.
Cream of Tartar 1.4 g.	No. 3
Venice Turpentine 16 g.	
Color 6-34 g.	Nickel Chloride 10 gr. Cobalt Chloride 10 gr.
00.00	
Tartar and soda are first dissolved in	
glycerin.	Dissolve. The writing becomes green
	on heating.
Varnish for Lithographic Inks	No. 4 Lead Acetate 10 dr.
Sandarac 15 kg.	Distilled Water 1 oz.
Olive Oil 15 kg.	
White Beeswax 12.5 kg.	Dissolve. The writing is invisible, and
Stearic Acid 12.5 kg.	becomes black when damped with a sul-
Oleic Acid 2.5 kg.	phide solution.
Castile Soap 2.5 kg.	70101 1 01 11
Burgundy Pitch 40 kg.	Billiard Chalk
Stearin Pitch 10-20 kg.	Formula No. 1
	Calcium Carbonate, Pre-
77 11 0 Audicale Tolicate	cipitated 115 g.
Varnish for Artistic Prints	a. Gypsum, Calcined 35 g.
Medium Strength	Pigment Powder (Blue,
Colophony, Pale 110 g.	(Green) 50 g.
Copaiba Balsam 70 g.	b. Borax Water (2%)
Tolu Balsam 2.5 g.	about 180-200 g.
Benzoin Amygdaloid 3 g. Lingued Oil 50 g.	to make a pasty liquid
	This paste is poured into slightly oiled
Dissolve hot.	molds.
	No. 2
30.31 Yamish /dan Tub-1	Calcium Carbonate 100 g.
Medium Varnish (for Inks)	Gypsum 30 g.
Rosin Oil 50 g.	Borax Water (2%) 115-130 g.
Sulphonated Rosin Oil Soap 3.5 g.	As above.
Boiled Weak Linseed Oil 4 g.	110 010 TO
Boiled "Middle" Linseed	Cellulose Tranfer Inks
Oil 52 g.	Formula No. 1
Colophony 25 g.	
By removing the weak linseed oil, a strong varnish is obtained.	Cellulose Acetate 170 os.
strong varnish is obtained.	Triacetin 200 oz.

INKS A	ND MARE	CING COMPOUNDS	193
- 1 Di I Berin	200 oz.	Printing Roller Cle	nnar.
High Phenol Resin	250 oz.	,	
Pigment	200 024	High Test Benzine	90 fl. oz. 10 fl. oz.
No. 2		Petroleum	10 IL 02.
Nitrocellulose (½ sec.)	15 oz.		
Triphenyl Phosphate	20 oz.	General Printing Cl	
Blown Castor Oil	5 oz.	High Test Benzine	80 fl. oz.
Basic Dye	2 oz. 50 oz.	Xyleno	15 fl. oz.
Acetone No. 3	JU 02.	Petroleum	5 fl. oz.
Nitrocellulose (1/2 sec.)	15 oz.	Intaglio Printing Pres	s Cleaner
Glyptal Balsam	20 oz. 5 oz.	High Test Benzine	80 fl. oz.
Stearic Acid	10 oz.	Tetralin	20 fl. oz.
Pigment Acetone	50 oz.		
No. 4	•	Off-Set Printing Cl	eaner
7 7	15 oz.	Use light petrol (gasoline	
Nitrocellulose (½ sec.) Phenol Formaldehyde Resin		Cho ngao peorer (ganorias	,.
Beeswax	50 oz.	Ink Remover	
Acetone	50 oz.	1	
No. 5		U. S. Patent 1,968	3,304
	50 oz.	A substantially non-aqueo	ous cream for
Triphenyl Phosphate Butyl Tartrate	50 oz.	the removal of ink stains i	from the akin
Cellulose Acetate	50 oz.	containing about 500 g. of	zinc stearate,
Mineral Oil	5 oz.	about 300 g. of citric acid,	about 500 cc.
Basic Dye	20 oz.	of 95% ethyl alcohol and a	bout 2000 ec.
No. 6		of diethylene glycol.	
Ethyl Cellulose, High		7.1 P. Norton	
Viscosity	50 oz.	Ink Eradicator	
Castor Oil	25 oz.	Potassium Alum	2 lb.
Mineral Oil	10 oz.	Citric Acid	2 lb.
Bronze Powder	20 oz. 50 oz.	Mix thoroughly and disso	
Benzol	50 02.	Water	3 lb.
Emulsifiable Transfer	Ink	Stencil Conting P	
Diglycol Stearate	20 oz.	U. S. Patent 2,011	1,898
Ethyl Cellulose	5 oz.	Formula No. 1	l
Sodium Abietate	10 oz.	Calcium Olcate So	lution
Pigment	10 oz.	Calcium Oleate	20 oz.
		Mineral Spirits	80 oz.
Ink Remover		The above ingredients are	e combined by
For cleaning dry printing	ink from	heating for a short time	in a steam-
printers' rolls and type.		jacket kettle.	
Denatured Alcohol	21/2 gal.	No. 2	
Commercial Toluol	14 gal.	Ammonium Stearate Soli	ution
Heavy Naphtha	3% qt.	Ammonium Hydroxide	14
Creosote Oil	1¼ gal.	(sp. gr. 0.9)	0.41 ox.
		Water	98.84 oz.
		Stearic Acid	0.75 oz.
Non-Inflammable Ink Re		The stearic acid is br	oken up into
(for Washing Printers' Rolls		small pieces and agitated	MILE THE OTHER
Carbon Tetrachloride	10 pt.	ingredients until dissolved.	* *
Toluol	13 pt.	No. 3	
Heavy Naphtha	11 pt.	Ammonium Oleate 8	Solution
Creosote	2 pt.	Ammonium Hydroxide	
		(sp. gr. 0.9)	0.41 oz.
Printing Form Clean	er	Water	98.84 oz.
Use light gasoline.		Oleic Acid	0.75 oz.

The above ingredients are combined in the same way as those of No. 2.

Suitable compositions for steneil paste in which the false bodying agents are incorporated are given below. The composition of the particular resin used is given after the examples setting forth the steneil paste compositions.

No. 4

white Stencii Paste	J	
Lithopone	46.1	oz.
Zinc Öxide	23.1	
*Resin A	15.7	oz.
Drier	1.5	oz.
Ammonium Stearate Solu-		
tion of No. 2	2.3	oz.
Calcium Oleate Solution		
of No. 1	4.8	
Mineral Spirits	6.5	oz.

No. 5

The same composition as No. 4 except that 23 parts of the lithopone are replaced by 23 parts of duatomaceous earth. The effect of the soap solutions described in the preceding examples is enhanced by the use of cellular or fibrous materials such as diatomaceous earth or "Asbestine."

No.

The same composition as No. 4 except that basic lead carbonate is substituted for lithopone.

No.

The same composition as No. 4 except that resin B is used instead of resin Λ .

No. 8

Black Stencil Paste

Differ During Lasto		
Carbon Black	17	oz.
"Asbestine"	4.1	()Z.
†Resin B	64	oz.
Drier	4.1	oz.
Ammonium Oleate Solu-		
tion of No. 3	7.2	oz.
Calcium Oleate Solution		
of No. 1	3.6	0Z.

No. 9

The same composition as No. 8 except that 7.2 parts of ammonium cleate solution are replaced by 4 parts of mineral spirits and 3.2 parts of calcium cleate solution of No. 1.

No. 10

Red Stencil Paste

Toluidine Red	19.8 oz.
Barytes *Resin A	28.6 oz.
*Resin A	21.8 oz.
Ammonium Stearate Solu-	
tion of No. 2	15.4 oz.
Mineral Spirits	12.8 oz.
Drier	1.6 oz.

The linseed oil modified resin given in this formula may, if destred, he replaced by a resin modified by linseed oil acids such as indicated by resin C below.

The ingredients in the pastes described above are combined in accordance with the usual products of paint manufac-

The following resins are illustrative of the class of polyhydric alcohol-polyhads acid resins especially suitable for the purposes of the present invention. These resins are made in the conventional way by reacting the ingredients in the proportions indicated.

*Resin A

12.8 oz.

27.1 oz.

55.8 oz.

Glycerol

Phthalic Anhydride

Linsced Oil Acids

Phthalic Anhydride	28	
Linseed Oil	59.2	0 Z .
†Resin B		
Glycerol	15	0 Z.
Phthalic Anhydride	35	oz.
Linseed Oil	50	0 Z.
Resin C		
Glycerol	17.1	0 Z .

LEATHER, SKINS, FURS

Chamois Leather from Rejected Calf Skins

pasted with The skins are soaked. sodium sulphide 1 and calcium oxide 4 (25° Bé.) at a temperature not exceeding 30° C., limed with calcium oxide 10 g. per liter, sodium sulpliide 4 g. per liter, water 400%, at 20° during 18-20 hours, washed with water at 22° for 40 minutes, fleshed, treated with 0.3% hydrochloric acid and 2% sodium chloride (of the weight of the raw skins) at 25°, softened with a concentrated softener (0.1% of the weight of the raw hide), for 1 hour at 35-37°, pickled for 40 minutes with hydrochloric acid 1.7, sodium chloride 7 and water 80%, tanned with chrome extract of 2% chromic oxide, having a basicity of 50%, split, neutralized, washed, greased, with 0.5% alizarin oil, 2% egg yolk and 150% water, washed the 20% of the chrome extract of the chrome oxide the contract of the chrome extract of the chrome ex with water at 35°, dried at 35°, let stand 2 days, dehaired in sawdust, stretched, cut, sand-papered and soaked.

Chamois Leather of Natural Color from Rejected Kid Skins

The skins are soaked in water at 18-20° C., drummed for 45 minutes at 17°, fleshed, soaked again in water at 16-17° drained and treated with a mixture of sodium sulphide 2%, calcium oxide 5% (of the soaked skins) of 30° Bé. at a temperature of 35-40°. The hair is removed by hand and the skins are placed in a lime solution for 5 days at 12-16°. They are then washed for 30 minutes, split, the thin parts are tanned by the formalin-fatty method and the heavier parts are chrome-tanned. The flesh side is treated with 0.5% hydrochloric acid for 45 minutes. The skins are further pressed and drummed in 5% of, seal fat, and treated in an oxidizing chamber for 14 hour at 32-35°. The above processes are repeated except that the oxidizing drying is carried out at 40-42°. The product is stored for 3 days, degreased with 200% water at 45° and sodium carbonate solution (5% of the weight of the skins) is added, the liquid discharged and the above soda solution again added together with water. The goods are sonked with water at 40-45°, drained, dried and stretched. They are dyed with nigrosine, drimmed for 6-7 minutes and fit liquored with 0.75% castor oil, 2% alizain soap and 2% rosin soap.

Velure from Rejected Pig Skins

Soak the raw hide in pieces weighing 1-3.5 kg. to a liquid factor of 1.5, at 20° C. and for 2 hours trent with 1 part sodium sulphide and 3 parts calcium oxide, density 25° Bé., at 25° let stand for 3-6 hours, unhair, wash and sort. Then treat with sodium sulphide 10 g, and calcium oxide 10 g. per liter at 20° for 4 days, split to an average thickness of 1.25 mm. and wash to a liquid factor 1:5 at 20° for 2 hours. Treat with concentrated softener 0.5% at 37° for 2-3 hours, de-ash with hisulphite 2% at 28° at a liquid factor 1:5; wash to a liquid factor of 1:5 at 25° and during 30 minutes. Pickle with sulphuric acid 2%, sedium chloride 10% and water 80% for 40-60 minutes at 18°. Tan with chrome extract containing chromic oxide 1.8, basicity 45 and water 80%; to complete tanning the basicity may be raised if necessary. Neutralize with bicarbonate 1.25 and water 200% at 35°, wash with water 300% at 40° for 30 minutes. Fat liquor with alizarin oil 1, egg yolk 3, water 150% at 40° for 40 minutes, and wash with water 300% at 35° for 25 minutes. Dry at 35°, unhair in sawdust containing 60% water for 16-20 hours, stretch, cut and polish.

Chrome-Tanned Black Calf-Leather

A calf leather which was previously tanned is planed on the grain side, neutralized, treated with 2% of pure fats, dried, unhaired and nailed on frames. The skin is then worked over: with grinding stones and the final treatment is given with pumice stone. Skins with a light nap are worked over with a wire brush (by hand). The skins are finally dyed with 15% (of their dry wt.) of substantive dyes and 4.5% ammonia, the mixture being diluted with 50% water.

Preparing Leather from the Mucous Stomach Membrane of Cattle

(1) The material is soaked, slightly fleshed, limed for 2 days, with about 12% slaked lime on the weight of the tissue, washed and delimed with bisul-phite. Tanning by vegetable or by onevat or two-vat chrome methods is followed by the usual dyeing, fat liquoring, drying and finishing. (2) The membrane is soaked for 2 hours in cold water, then for 15-20 minutes each in 3 vats with a gradually increasing temperature from 22° C.

Removing Scales from Shark Skins

Give the skins a salting in a 1% solution of sodium chloride. Then a treatment in a 1/2% solution of hydrochloric acid. This method should dissolve the scales, but if for any reason it does not, keep on increasing the percentages of both materials. Then give the skins a thorough washing in pure water in a Watch carefully that the hydrodrum. Watch carefully that the hydro-chloric acid does not attack the skins themselves.

Loosening Hair from Hides Canadian Patent 353,326

Wheat Shorts 14 lb. Wheat Bran lb. 0.6 cc. Phenol Solution (21/2%) Water 15 gal.

Preparing Pigskins for Tanning

First, scrape the raw skins until they are nearly dry. Then give them a good soaking for a day or two. Next wash Next wash them in a drum or vat containing a warm solution of sal soda or similar product for loosening the grease. In preparing this solution, use from 1% to 2% of sal soda according to the condition of the skins, i.e., they appear to be extremely greasy, a higher percentage of sal soda is preferred. After the skins have reseived a thorough soaking in this solution, strike them out thoroughly with a dull knife, forcing out as much grease as possible. Very greasy skins should be struck out two or three times. Then rinse them off in warm water and soak them overnight in cold water, after which they are unhaired and limed.

As pigskins absorb tan liquors some-what slower than calf and other skins, it is good practice either to give them slightly stronger liquors or a longer time in the same strength liquors you are Water 96.5 lb. using for your other stock. This sugges- for 30 minutes, moving repeatedly.

tion applies more especially to a vegetable tannage.

Pigskins being of a very greasy nature require less oiling or fat liquoring than other skins. Some tanners reduce the oiling from 20% to 30%.

Felting Animal Hair German Patent 608,770

Hair is rendered capable of fulling and felting by treatment with a bath containing small amounts of oxy acids of metals of the chromium group or their salts together with hypochlorous acid or persulphuric acid or their salts. Thus pelts are treated with an aqueous solution containing 2% potassium chlorate, 1% nitric acid and 0.1% chromium in the form of dichromate at 10-100° C., and dried.

Treating Lizard Skins

Bleaching should be effected in two solutions. (1) potassium permanganate 5 g. per liter, sulphuric acid 1 g., water 500% of the weight of the skins, and (2) water 500%, bisulphite 25 g. per liter. The washed skins are dyed beige by treating with 0.03% orange PB, 0.04% methanyl yellow and 200% water for 20 minutes, adding 0.3% acetic acid and treating 20 minutes. For gray use nigrosine 0.1%, acid brown 0.01% and water 200% at 45° for 15 minutes; add 0.3% acetic acid and treat for 15 minutes. For violet use wool brown 0.5% and acetic acid 0.5% at 45°, add 0.1% methyl violet after 30 minutes and treat for 15 minutes. For blue use sulphone acid-blue 0.3% and water 200% at 45° C. for 15 minutes, add 15% acetic acid and treat for 20 minutes.

Bleaching Deer Skin

Formula No. 1

Make a bath with

Hydrogen Peroxide (30%) 5-8 0.5 lb. Seignette Salt

and put the skins into it for 1/2 hour. Dry them thereafter at 30° C. If the skins are not pale enough, repeat in the same bath.

No. 2

Put skins into a solution of Potassium Permanganate Sulphuric Acid 0:5 lb. 96.5 lb. Water

5 lb.

95 lb.

Wash out in cold water, then in solutions of

Sodium Bisulphito Powder 5 lb. Water 95 lb. (for ½ a minute), and

Hydrochloric Acid Water

(for 1/2 minute).

Then wash out very carefully, repeat the process until the wanted paleness is reached.

No. 3

Tanning After Bleaching (Often Advisable)

Wash for 2 hours at 30-35° C. in solution of sodium carbonate, spill with water, and treat for 7 hours in a solution of

Sodium Carbonate 2 lb. Formaldehyde (40%) 2.5 lb. Water 95.5 lb.

Tanning Greenland Scal Skins

The sorted skins are soaked in water for 10 minutes, fat is removed from the flesh side and the skins are again soaked in water for 36 hours with change of They are water at 12-hour intervals. degreased in a drum charged with water of 30° C. with addition of 1% sodium hydroxide (calcined on the salted skins). The skins are washed in running water for 30 minutes, drained on racks for 2 hours, placed for 30 minutes at 25-30° in a solution prepared from sodium sulphide 20 g. per liter and calcium hydroxide 160 g. per liter, unhaired with a tool, washed till the concentration of sodium sulphide amounts to 20-25 g. per liter, and treated for 2 days in a lime solution used once for unhairing, with addition to the solution of 12 g. calcium hydroxide in the course of the processing. The skins are then washed, split, delimed and tanned in a six vat battery for 6 days, with a 4° Bé. solu-tion in the first and 4.25° Bé. in the last vat. The drum tanning may be carried out in an oak extract of 7° Bé. The aging in stacks requires 24 hours and the deacidification, which is carried out with 1° Bé. solution during 4 hours, is followed by washing in running water for 8-10 hours.

Tanning Horsehide
Full Grain Horse for Glove and Sport

Having selected hides after unhairing for this type of leather, they are pickled,

tanned, pressed, staked, etc., in the same in anner as buffed glove horse. The stock is then splt and shaved. After this it is neutralized and fat liquored in the same manner as for the "One Bath" tanned stock which is given below.

One Bath Tannage for Full Grain Horse

Often a tanner prefers to tan glove horse leather with the single bath tannage rather than with the two bath tannage. The final results will be the same, as both tannages produce excellent leather.

After lime splitting, the stock is bated and washed and taken to the chrome tan wheels. Maximum loads of 3000 lb. of lime split stock are considered sufficient.

The taumage is based on this weight.
Place the stock into the drum with 180
gal, of water and 180 lb, of salt, mill for
10 minutes, then add 45 lb, 66° 136 sulphanic acid in 15 gal, of water. Mill for
55 minutes, then add 42 gal, of chrome
liquor.* This is added in three doses of
14 gal, each, 30 minutes spart. After
the last addition is made, continue milling for 4 hours, let stand in drum over
might.

The following morning mill the stock 1/2 hour, then add:

Fifteen pounds bicarbonate of sods, first dissolved in 20 gal, of water. Add thus at the rate of 1 gal, every 2 minutes; continue milling 30 minutes, remove from the drum, by flat on trucks, let drain for 24 hours, set out, split and shave.

* The chrome liquor used for this purpose is made as follows:

Use a lead lined tank. Place the bichromate, alumnum sulphate and 200 gallons of water into the tank, agitate well by means of an air line, then add the sulphuric acid. The corn sugar is made to a syrup with water and is added very slowly, taking the usual

and is acted very stown, states and all acted very stown, so the sugar has been added, add five gailons of bisulphite of sods, 33° Baums, boil the liquor for one-half hour, allow to cool and make up to 500 gailons, stir wall and allow to age ten days before using.

Coloring

Divide the tanned split stock into lots of 400 lb. each for coloring and fat liquoring. Place the stock into the drum with 120 gal. of water at 90° F., then add 6 lb. of bicarbonate of sods dissolved in 20 gal. of water, and mill for ½ hour. Drain the drum and wash the stock for 1 hour at 80° F., again drain

the drum and add 200 gal. of water at 120° F.

Prepare the following dye mixture:

Fustic Crystals 2 lb.
Resorcin Brown 4½ oz.
Fast Red ½ oz.

Boil together in 30 gal. of water, cool to 125° F., and add to the drum. Mill stock in the dye solution for ½ hour, then drain the drum.

This will produce a cream color which is a standard for glove and sport goods stocks. The amount and type of fat liquor determine the purpose to which the stock will be used.

Fat Liquor for Stretchy Glove Leather
Sulphonated Cod Oil 24 lb.
Sulphonated Mineral Oil 24 lb.
Sod Oil 24 lb.
Borax 4 lb.

Place the materials into a barrel in the order given, stirring well upon addition of each item. Add 25 gal. water and heat by means of a steam jet agitator to 160 F. The total volume should be 50 gal. This emulsion is added to the drum after draining off the exhausted dye solution. Mill for one hour, rinse very slightly with water at 100° F., take out of drum and horse up for 24 hours, then hang up to dry.

Fat Liquor for Sporting Goods Leather Sulphonated Mineral Oil 64 lb. Sod Oil 24 lb. Borax 4 lb.

Place the materials into a barrel in the order given, stirring well. Then add 25 gal. of water and heat by means of a steam jet agitator to 160° F. The total volume should be 50 gal. This emulsion is added to the drum after draining off the exhausted dye solution. Mill for 1 hour, rinse slightly with water at 100° F., take out of drum, horse up 24 hours, then hang up to dry.

Drying

This type of leather can be dried rapidly. Since it is quite wet, the initial air temperature can be 120 to 130° F. Rapid circulation of the air must accompany the high temperature; the moist atmosphere is gradually expelled from the dry room, emitting at the same time fresh and reducing the temperature, so that the stock is thoroughly dried in 24 hours.

Crusted Stock

After the stock is dry it is crusted for five days. Dip the crusted stock in

water at 110° F. for one minute, place into bins, cover well with damp burlap and allow to mull for four hours. Then place into damp sawdust (containing about 35% moisture) and let it rest for 24 hours. Then stake on a Slocum Machine and hang up to air off for an hour.

Dry Mill

After the stock has aired, place into a dry mill. For each 100 sides use 10 to 20 lb. of French chalk, the amount depending upon the size of the stock. Dry mill for 1 hour. Remove from dry mill and stake on the Baker Machine. After the second staking, polish the grain on a shearling wheel.

Notes: Some adjustments may have to be made for either the "one bath" or the "two hath" operations. In a greater number of cases the adjustment is made in the fat liquor stage, either increasing or decreasing the amount. Drying of the stock must be carefully controlled since this operation is very important to a soft, yet full feeling leather.

Leather of this type should not be tacked. Leather of this type should be stretchy, the glove more so than the sport leather. The latter is used principally for baseball gloves.

Black Garment Horse Leather

This type of leather is used principally for coat stock, although it can also be used for glove purposes. The market for this leather is highly competitive and therefore the leather must be made as economically as possible. Sheep, in grain and suede, is used very extensively and is produced at a low cost including the raw material. Because of this, it has found a greater market than horse leather. For general utility and durability, horse garment leather excels sheep leather.

The stock is sorted in the beamhouse before bating. The butts should be split down to a minimum. After bating and washing, the stock is transferred to the chrome tan yard.

A maximum drumload of 3000 lb. of lime split stock will be used. The stock is placed into the drum with 200 gal. of water at 65° F. and 180 lb. of sait. Mill 5 minutes and then add 42 lb. sulphuric acid, 66° Bel, in 15 gal. of water, and mill 15 minutes, then add 45 gal. of chrome liquor.* This is added in three doses of 15 gal. each, 30 minutes apart. After the last addition of chrome liquor mill for 5 hours, let stand in drum overnight.

The following morning, mill the stock for 30 minutes, then add 15 lb. of bicarbonate of soda dissolved in 20 gal. of water at 75° F.

Add the soda at the rate of 1 gal. every 2 minutes. After the last addition mill the stock for 30 minutes, remove from drum and horse up for 24 hours, set and split. The split stock is divided into lots of 500 lb. each for coloring and fat liquoring.

* The chrome liquor for this tannage is made as follows:

auc as tonows.	
Bichromate of Soda	1000 lb.
Sulphuric Acid, 66° B6.	980 lb.
Corn Sugar	832 lb
Total Volume	500 gal,

The usual precautions must be taken and the manner for procedure is the same as that for the chrome liquor under "One Bath Tannage" for glove horse.

Coloring

Place the stock into the drum with 150 gal. of water at 90° F., and add 334 lb. soda ash in 10 gal. of water. Mill for 30 minutes, then wash with water at 110° F. for 1 hour. Drain the drum and add:

Water at 120° F. 250 gal. Direct Black in 30 gal. of Water at 120° F. 17½ lb.

Mill 30 minutes and drain the drum, add:

Water at 120° Methyl Violet	F.	250 24	gal.
and Acetic Acid		4	02.
in 20 gal. of Drain drum,	water,	mill 20	minutes

Fat Liquoring

Prepare the following:	•	
Logwood Crystals	71/2	lb.
Water, Boil and Add	20	gal.
Fig Soap	15	lb.
Sod Oil	100	lb.
Sulphonated Cod Oil	10	lb.
Total Volume	50	gal.
True of the teacher		-

Use steam jet agitator for the purpose of preparing the above emulsion, add to the drum at 150° F., and mill 1 hour. Remove from drum and horse up to drain for 16 to 24 hours, set out on Turner Serial Table Machine.

Oiling and Drying

Oil off the set out stock on the grain with a light paraffin oil, using a shearling swab for the purpose. Apply a light coat. Then send the stock to the dry room. Hang up the stock in a room equipped with fans and heating coils. A temperature of from 90 to 100° F. is maintained; the air is well circulated with fans so that drying is effected in

24 to 36 hours. The stock is then crusted for two days in a cool room.

Sammying and Staking

Dip the stock in warm water, 110° F., for I minute, place into a bin, cover with burlap and allow to mull for 4 hours, then place into damp sawdust containing 40% moisture, let rest for 24 hours. Then stake on a Slocum Machine equipped with a fiber pad on the staking head. Apply as much pressure as the stock will stand; cracking of the grain must be avoided. Then hang up the stock to air off at room temperature, restake and trim closely where necessary and again stake if hard spots are found.

Finishing

Use the following finish:

cae the following maisn;	
Shellac Solution	6 pt.
Casein Solution	814 pt.
Liquefied Gelatin	635 pt.
Carnanba Wax Emulsion	11/2 pt.
Sulphonated Cod Oil	1 pt.
Nigrosine	1¼ lb.
Water	30 pt.
Ammonia	1 pt.

Mix the above ingredients in the order given, the Nigrosine first dissolved in the water. Apply two conts of the finish to the stock, allowing to dry well after each application. Finally polish on a shearling wheel.

In order to obtain the desired results it may be necessary to vary the quantities of some of the finish materials. A third coat of finish may also be required. Proper drying between coats is of importance.

The greatest factor affecting finishing of leather is the type and amount of fat-liquor used. This holds particularly when a finish job at low cost is desired. In other words, the finish must be properly adjusted by varying its components until the proper balance is obtained.

Synthetic Tanning Process U. S. Patent 1,975,616

The hides, skins or pelts are prepared by any suitable and well known process and then immersed in a solution containing approximately 20% of a urea-formal-dehyde solution and 10% of salt at about 35° C. and gently agitated for about 5 hours. The temperature may then be raised to 45° C. and the solution acidified to about pH₃ with sulphuric acid and agitation continued for 30 minutes. The temperature is then raised to 55° C., the skins worked for 15 minutes, cooled, rinsed in cold water, neutralized with

200 THE CHEMIC	AL FORMULARY
sodium bicarbonate, rinsed, fat liquore	d Leather Fat, Yellow
and dried.	1
One method of producing the ures	Formula No. 1 No. 2
formaldehyde solution mentioned in th	e Parama wax 8.5 5 kg.
above example is as follows: 3 oz. o	
urea, 1½ oz. formaldehyde, 2 oz. sodiur	
carbonate and 16 oz. of sodium chlorid	
are dissolved per gallon of water, an this solution employed in the tanning	Yellow 1435, Dye 10 10 kg. Carnauba Wax — 1 kg.
process at once, or at least prior to th	Wool Fat - 0.3 kg.
formation of an insoluble precipitate.	
	No. 3
Leather Oil	Paraffin Wax 8,000 g.
	Carnauba Wax 1,375 g.
Spindle Oil 96 g. Caoutchouc, Crude 3 g.	Wool Fat 340 g.
Caoutehouc, Crude 3 g. Resin, Coumarin, Viscous,	Train Oil 5,670 g. Mineral Oil 35,000 g.
Liquid 1 g.	
Heat to 100° C. and stir until dis	Yellow 1435, Dye 12 g.
solved; add a little Birch Tar Oil (a	No. 4
perfume).	Paraffin 10,000 g.
St. of St	Ceresin 9,000 g.
Sport Leather Oil	Carnauba Wax Arrears 1,000 g.
Pale Train Oil 50 g.	Train Oil 4,000 g.
Degras 20 g.	Spindle Oil 70,-80,000 g. Yellow 1435 20 g.
Woolfat, Neutral 5 g.	Yellow 1435 20 g.
Birch Tar Oil 5 g.	
Spindle Oil, Refined 20 g.	7
Melt together and add:	Leather Dressings or Finishes
Caoutchone Solution (5-10%)	Formula No. 1
in Toluol 2 g.	Shellac 9 g.
	Venetian Turpentine 1 g.
Oiling Leather with Petrolatum	Castor Oil 1 g.
Satisfactory penetration is obtained by	Alcohol 89 g.
drumming with a hot mixture of petrola	Mix until dissolved and filter.
tum 45, mineral oil 40, and degras 15%.	No. 2
Special Leather Oil	
Cold Test (20°) Neats-	Gum Sandarac 5 g. Castor Oil 2 g.
foot Oil 50 gal.	Alcohol 81 g.
Paraffin Oil (28°) 25 gal.	No. 3
Water 25 gal. Sulphonated Castor Oil	0 (1) 11
(50%) 25 lb.	Orange Shellac 16 g. Caustic Soda 0.9 g.
Manipulation: Mix water first with the	
sulphonated castor oil. Then mix all in	
gredients at 30° C.	Water 81 g.
-	No. 4
Leather Fat, Black	Orange Shellac 27 g.
Formula No. 1 No. 2 No. 3	Caustic Soda 214 g.
	Boric Acid 24 g.
777	Sodium Ricinoleate 214 g.
Wool Fat, Raw 2 1 — g. Montan Wax,	water 60 % g.
Crude 4 3 3.9 g.	No. 5
	Shellac, Bleached 20 g.
Carnauba Wax,	
Gray 2 — g.	Galipot Resin ½ g.
Gray 2 — g. Nigrosine, Fat-	Borax 4 g.
Gray 2 — g. Nigrosine, Fat- Soluble 1 0.3 0.39 g.	Borax 4 g. Ammonium Hydroxide ½ g.
Gray 2 — g. Nigrosine, Fat-	Borax 4 g.

5 g. 5 g. 2 g.

100 ğ.

100 g.

Dressing for Hunting Les	ither	
(Inflammable!)		
a. Nigrosine Base Olem	10 30	g. cc.
b. Benzol (90%) Alcohol Cleaning Benzoline	150 500 500	cc.
Auto Top and Artificial Leather Nitrocellulose (Film Scrap) Camphor Ethylacetanilide	40 10	essing g. g.

sing for Hunting Land

Suede and Chamois Leather Dressing U. S. Patent 2,015,943

Castor Oil

Lampblack Nigrosine

Alcohol

Benzol

Acetone	90 oz.
Chloroform	60 oz.
Liquid Petrolatum	140 oz.
Naphtha	870 oz.

Leather Finishes

A good polish is made from 22 g. stearin, 22 g. carnauba wax and 56 g. linseed oil. It is better to prepare an "emulsion polish" by mixing 22 g. stearin, 22 g. carnauba, 11 g. paraffin, 23 g. linseed oil, 3 g. ammonium chloride and 17 g. water. The carnauba wax may be replaced by synthetic waxes. Waterproof spirit finishes are made by mixing shellae (9 g.), Venetian turpentine (1 g.), castor oil (1 g.), and 96% alcohol (89 g.); or mastic (12 g.), sandarac (5 g.), castor oil (2 g.), and spirit (81 g.). All grease should be removed from the leather before application of spirit finishes. For making polishes of good elasticity a recipe recommended is: ruby shellac (16 g.), technical caustic potash (0.9 g.), boric acid (1.2 g.), castor oil soap (0.9 g.), water (81 g.). Camphor oil may be added as a perfume. For treating leather of more porous nature, colloidal matter such as carragheen moss, algin, etc., are added to the above soap finishes, or gum tragacanth may be used. A recipe for green bronze finish is magenta 7.6 g., safranine 1.9 g., ruby shellac 1.4 g., and methanol 89.1 g.

Fur Glazing

Dissolve 3 to 6 oz. of paraffin wax in 1 gal. petroleum cleaning solvent.

Approved cleaning solvent is preferable because of its safety during ordinary handling.

Precaution: Paraffin separates from the petroleum solvent at temperatures below 70° F. At --15° F. it is completely chilled out of the solvent.

This finish is used for the saturation of dry cleaned furs to replace any oils removed and to make them water repellent. It is also sponged or sprayed on materials that are lifeless or lusterless after cleaning and drying to produce high gloss.

Natural Color and Glaze for Snakeskins An alum tannage is good for pocketbook leather and will as a rule impart a natural color. For each 100 lb. of bated and drained skins use 7 lb. alum, 2 lb. salt, 8 lb. flour and 5 lb. liquid egg yolk. The alum and salt are first dissolved in a small quantity of hot water and the solution then cooled. After cooling, the solution is added to the flour with constant stirring. Dissolve the egg yolk separately in a small quantity of cool water and then add to the other ingredients. This mixture when ready to apply should weigh about the same as the skins, that is, it should measure about 10 gal. for every 100 lb. of skins.

Stir the skins in this mixture for about 3 hours, or until nearly all of it is absorbed by the skins. Leave the skins in the same container or vat overnight. Then strike them out, stretch moderately on boards and dry. After drying, take the skins off the boards and wash them with a brush in cool water. This washing will remove any dried mixture remaining on the grain side.

maning on the grain side.

Next lay the skins in piles overnight with grain to grain and cover with a moist cloth. Then stake and dry. After drying give the skins another staking. Some tanners also fluff the flesh side.

A good mixture for glazing can be made from the following: 1 oz. egg albumén, ½0 oz. gelatin, 2 oz. milk and 5 pt. water. The egg albumen is dissolved in 4 pt. of the water at 90-95° F. and the gelatin dissolved in 1 pt. of hot water and then allowed to cool to 90-95° F. The two solutions are mixed and then the milk is added.

This mixture is brushed on the grain side. The skins are then dried again and glazed by machine. Some tanners repeat this application and add a small quantity of casein or shellac. Others use castor oil and methylic alcohol.

Dressing Bagdad Leather

Skins known commercially as Bugdads differ considerably in weight, size and quality, but they are all usually heavily loaded with dirt and loose tanning matters, all of which require to be completely removed before the goods can be properly dressed. After sorting, trimming and perhaps necking on the shaving machine, the goods need drumming for half an hour in a solution made up of 10% salt and 14% sulphuric acid on the dry weight. Some tanners use a cold solution, but a temperature of 100° F. will be found advisable for complete action. The object of processing the goods in the above liquor is to cleanse and open the pores of the leather so that it will be able to absorb the tannins during the next stage of dressing. At the end of the allotted time, namely half an hour, the liquor should be run off and the goods washed up in running water, preferably warm, for three-quarters of an hour. If the Bagdads are in a filthy condition the percentage of sulphuric acid should be increased to 1%, and this will generally prove strong enough to clear the grain and remove any stains, par-ticularly iron marks. These preliminary processes are very important, especially in the case of whites, where it is of the utmost importance that the leather should be as clean as possible before the bleaching or whitening process commences.

Re-tanning

This operation can be successfully carried out in the drum, and, indeed, this is really the most suitable receptacle. A good synthetic tanning material, such as Maxyntan or Sellatan, in conjunction with sumac extract, usually forms the basis for a white tannage, and it is not advisable to use any tannin likely to darken the color of the leather. A run for half an hour in 5% of the synthetic followed by half an hour in 5% sumac extract will be found eminently satisfactory, but if it is necessary to reduce expenses to a minimum, the synthetic can be increased and less sumac extract employed. The tannage gives a very clean and fairly soft leather which will feed up well. The amount of water used depends a great deal on the weight and size of the goods, but in all cases the minimum should be run in, as this will ensure better exhaustion of the liquor.

After re-tanning for one hour, the goods should be taken out of the drum and horsed up overnight. Whilst this is not absolutely necessary, it is always ad-

visable if time and labor charges will permit, as it enables the tan to fix and the fibers to feed. Practical experiments have shown that there is a recognizable difference in the handle of leather allowed to drain for 12 hours as compared with leather rushed through the processes.

Bleaching

Next day, run the goods in the following solution: 214% barium chloride and just sufficient warm water, 100° F., to cover the leather.

A run of a quarter of an hour will enable the leather to take up the barium salt and exhaust the solution. An addition of sodium sulphate, 5%, dissolved in a small volume of warm water will precipitate barium sulphate, a white insoluble salt, in the fibers of the leather. This bleaching process is quite economical and if worked properly it will be found to give a very clean, white leather.

Some tanners use sulphuric acid instead of the sodium salt, but sodium sulphato is equally satisfactory and with it there is less chance of the leather being rendered hard and brittle.

Whitening and Filling

To fill out the leather, improve its handle and general appearance, it is advisable to work the goods in the following mixture, which should be added to the drum through the hollow axle:

Devolite Clay	15	1b.	
Flour	15	lb.	
Soap	5	lb.	,
French Chalk	5	lb.	
Turkey Red Oil	21/2	lb.	
Trace of Methyl Violet.			

A run of three-quarters of an hour in this liquor will complete the operation and afterwards the goods should be horsed up for a few hours preparatory to striking out and straining. The former process must be well done in order to remove all the wrinkles and drawn grain. To retain the fullness and suppleness of a well-nourished leather, the latter should be dried out in a moderate temperature. It is a bad practice to dry the leather in a fierce temperature for the sake of a few hours, but if this is imperative, then the temperature should be increased gradually. When dry, the leather requires buffing, then chalking on the grain and flesh, and finally boarding.

Semi-chrome Colors

A better quality skin is unsally chosen for this work, and naturally the tanner has a better chance of producing a full and nice feeling leather. Goods should be washed in warm water for half an hour to remove loose dirt, and then stripped in a weak alkaline bath made up with 1 to 2% borax calculated on the dry weight of the leather. The stripping should take about an hour, and by this time practically all the loose tannin will be removed. The alkaline liquor should then be run off and the goods thoroughly washed in running water for half an hour.

Re-tanning in a Chrome Bath

After draining, the washed leather should be drummed with its own weight of a 4% salt solution for 10 minutes and the chrome liquor added. Prepare the chrome liquor by adding soda crystals to reduce the basicity. When using panchrome, 1 lb. of soda crystals for every 8 lb. of chromium salt is recommended. The latter should be dissolved in a known volume of hot water, and the soda dissolved in a small amount of hot water. The alkali must be added very slowly and the liquor stirred constantly during the addition.

The amount recommended for retanning Bagdads is 7% chromium salt on the dry weight of the leather. The chrome liquor should be passed into the drum through the hollow axle in three parts, at intervals of half an hour. A period of 2½ to 3 hours is recommended for complete re-tannage. The addition of 1% ordinary washing soda is then made, and drumming continued for a further hour. At the end of that time, the leather should be well tanned, and it is advisable to horse up for twelve hours or so. The next morning, the goods will need neutralizing, and 1% borax on the dry weight is recommended; a period of three-quarters of an hour will be found to be sufficient to neutralize the leather.

A light mordanting is recommended to ensure more level dyeing, and to give the leather a better feel or handle. Gambier is quite good, so also is Osage Orange Extract; about 2% on the dry weight will be found ample. Acid dyes should be used and there is, of course, an unlimited number of colors available.

After dyeing, the leather should be well fat liquored, and the following recipe is excellent for semi-chrome clothing leathers. Dissolve ½ oz. of potassium carbonate in a small quantity of hot water, 180° F., and then add 2 lb. of neatsfoot oil and ½ lb. of potash soap. Emulsify the mixture and then add 1 lb. of heavy sulphonated oil and ½ lb. of mineral oil and stir vigorously until the emulsion is stable. Use 4 lb. of this

fatty mixture for every 100 lb. of dry leather. After fat liquoring, the goods should be horsed up for several hours prior to striking out and drying. The drying should be carried out in a moderately warm, but not hot shed, and it is not advisable to have the goods strained, as it is likely to render the leather hard and impoverished.

When dry, the leather should be stored in damp sawdust for 12 hours or until in the right condition for staking. After staking and drying it requires fluffing on an emery wheel and finally dope finishing in the usual way.

Belt Dressing

Formula No. 1		
Wool Fat	50	ø.
Mineral Oil (0,885-90)	20	ø.
Paraffin Wax (56-58° C.)	10	
Ceresin, Yellow (58 60°)		ĝ.
Castor Oil ("Second Press-	-	Θ.
ing'')	10	ø.
Degras	5	g.
No. 2		ω.
Resin	40	ø.
Train Oil	10	ø.
Cotton Seed Oil or Sperm		ю.
Oil, Blown	15	g.
Paraffin Scale Wax		
(48-52° C.)	15	ø.
Mineral Oil (sp. gr. 0.905)	20	ġ.
No. 3		
[Wool Fat, Neutral	30	g.
a. Wool Fat, Neutral	20	g.
	10	
b. Graphite, Amorphous		-
Castor On	10	g.

Melt up the fats a, stir then into the fusion graphite, and castor oil. Press. The product is soft and like a salve.

Shoe Bottom Dressing

Montan Wax, Bleached	10	0 z.
Paraffin (or Scales), White (50-52° C.)		oz.
Anilin Dyestuff (Oil Soluble)	2	0 Z.
Turpentine Oil (or Sub- stitute)	54	0 z.

Patent Leather Dressing Black

Formula No. 1		
Celluloid	20	lb.
Castor Oil	5	lb.
Lampblack	5	lb.
Alcohol	30	lb.
Benzine	35	lb.

No. 2 Celluloid	25 lb.	No. 1601. The solvents are mixed and the dyestuff placed in a cloth sack and
Lampblack	8 lb.	suspended in the solvent mixture which
Nigrosine	1 lb.	is occasionally agitated.
Castor Oil	6 lb.	
Alcohol	20 lb.	
Benzine	45 lb.	Shoe Luster (Finish)
No. 3		Water 850 cc.
Celluloid	25 lb.	Ammonia (0.910) 20 cc.
Lampblack	8 lb.	Shellac, Bleached, Finely
Nigrosine	1 lb.	Powdered 150 g.
Castor Oil	8 lb.	Let stand cold for some hours; heat
Alcohol	25 lb.	the jelly formed to liquefy it.
Benzine	40 lb.	
Red	30 lb.	
Celluloid Ochre	5 lb.	High Luster Finish
Castor Oil	5 lb.	(Water 100 cc.
Zinc White	3 lb.	a. Borax 25 g.
Nigrosine	2 lb.	Shellac, Bleached 150 g.
Alcohol	20 lb.	b. Water 700 cc.
Benzine	30 lb.	c. Turkey Red Oil 50 cc.
Blue		Dissolve a, warming up gently without
Celluloid	30 lb.	boiling; thin with b, and add c.
Zinc White	5 lb.	g,
Paris Blue	2 lb.	
Castor Oil	8 lb.	Dark High Luster Finish
Alcohol	25 lb.	Ruby Shellac, Powder 150 g.
Benzine	25 lb.	Water, Cold 850 cc.
Green	00.11	Ammonia (0.910) 20 cc.
Celluloid	30 lb.	Soak for 6-8 hours (covered), warm
Zinc White	5 lb.	to complete solution (if necessary, add
Schweinfurth Green	2 lb.	more ammonia). Optional: add dyestuff.
Castor Oil	8 lb. 25 lb.	
Alcohol Benzine	25 lb.	
Denville	۵۵ ۱۵۰	High Luster Finish
		(Ruby Shellac 150 g.
White Shoe Bottom	F 'inish	a. { Water 200 cc.
Gum Tragacanth	2 oz.	Ammonia (0.910) 30 cc.
Water	1¼ gal.	b. Water 550 cc.
Soak and stir until smoot	h, then add	Make up a, thin with b.
Precipitated Calcium Car		ap w, onthe files of
bonate	2 lb.	**************************************
Titanium Dioxide	1/4 lb.	Liquid Burnishing Wax for Shoe Soles
Oxalic Acid	1 lb.	Carnauba Wax 20 oz.
Copper Sulphate	1 lb.	Turpentine 20 oz.
Magnesium Sulphate	5 lb.	Black Dye (Oil Soluble) 3 oz.
Sal Soda	3 oz.	Duponol W.E. or Lohrinol 5 oz.
Water	6 oal.	Forrig Acatata 6 or

Black Dye for Leather

Water

3 6 oz.

gal.

The following dye solution is used for the dyeing of the uppers of leather shoes. It will render same black in one applica-tion regardless of the previous color.

Black Dye (Alcohol Soluble) Methanol Benzol Nitrobenzol	66 2 0	0Z. 0Z. 0Z.
Nitrobenzoi	10	02.

The black dye should be of the acid type such as Calco Condensation Black

45.8 oz. Reduce the ferric acetate to a powder and dissolve same in the acetic acid and water mixture. Dissolve the Duponol W. E. in the above solution and heat to about 170° F. Melt the carnauba wax and pour into the turpentine which has been previously heated to about 180° F., dissolve the black dye in this mixture, and then add this latter solution to the former while agitating vigorously. Allow to cool with continued agitation. Du-

Ferric Acetate

Glacial Acetic Acid

в oz. 0.2 oz.

ponol W. E. is one of a series of soaps or emulsifying agents of the higher alcohol sulphates which are effective as such in	No. 3 Amber 380 g.
an acid solution.	Linseed Oil, Boiled 250 g. Sandarae 30 g.
	Turpentine, Venico 60 g.
Preserving Hides and Skins	Turpentine 200 g.
9	Tallow 600 g.
German Patent 617,166	Caoutchouc 75 g.
Salt 99 lb. Sodium Perborate 1 lb.	Linseed Oil 300 g.
Souther Ferborate 1 ib.	No. 4
	For Hunting Shoes
Conservation of Shoe Soles	Caoutchouc 4 g.
Melt up:	Pig Fat 6 g.
Linseed Oil 50-60 g.	Cod Liver Oil 24 g.
Paraffin 40-50 g.	No. 5
Heat 80° C.	For Horse Covers
Treat soles with this mixture after thorough cleaning, 2 or 3 times in 4-6 weeks.	Japanese Train Oil 94 g. Saturated Caoutchoue Solu-
WCCES.	tion in Turpentine 5 g.
77 1 A Cl C.l	And 1.5 g.
Hardener for Shoe Soles	
Rosin, Pale 4 g. Linseed Oil Varnish 5 g.	Quick Black Shoe Edge Ink
	Bright Drying Carnauba
Dissolve hot and add:	Wax Emulsion 50 lb.
Benzoline or Turpentine or Mixture 9 g.	Nigrosine 8 lb. Water 3 gal.
Mixture 5 g.	Water 3 gal.
Impregnation of Shoe Soles	
	Edge Filler for Shoe Factory Use
French Patent 750,728	Sonp 15 lb.
Benzoic Acid 3 g.	Yellow Dextrin 5½ lb.
a. Acetone 40 cc.	Neatsfoot Oil 1½ qt. Oil of Mirbane 1 pt.
-111001101	Gelatin 111/2 lb.
b. Oxalic Acid 3 g. 5 g.	Formaldehyde 1 qt.
Water 50 cc.	Water 1 qt.
Dissolve a and b separately, mix, add 15 g. of dye to 1 liter; brush on roughened soles.	This is made up with sufficient water to make 60 gal. solution.
D	Brown Shoe Heel Stain
Preservation and Hardening of Sole Leather	Alcohol 7 fl. oz.
Linseed Oil 6 cc.	Acetone 1 fl. oz.
Water Glass (40-45° Bé.) 4 cc.	Gum Tragacanth 4 oz.
Mix until emulsified. Apply with	Mix the above until gum is thoroughly
brush.	wetted and to it add slowly with stirring the following solution made by boiling
Waterproofing Leather	and then cooling:
Formula No. 1	Oxahe Acid 3 oz. Water Soluble Brown Dye 8 oz.
Gutta-Percha 2 g.	Water Soluble Brown Dye 6 02. Water 2½ gal.
Rape Seed Oil, Boiled 8 g. Yellow Wax 6 g.	Strain through cheesecloth.
Yellow Wax 6 g. Pig Fat 25 g.	Ditale through checkerous
Venetian Turpentine 60 g.	
Spermaceti 1 g.	Shoe Dye Remover
No. 2	Isopropyl Alcohol 7 cc.
Linseed Oil 100 g.	Acetone 1 cc.
Gutta-Percha 10 g.	Butyl Cellosolve 1 cc.
Copal Varnish a little	Water 10 cc.

Shoe Repairing Cement U. S. Patent 2,004,059

Six pounds crepe rubber, 2.5 lb. rosin, and 1.5 lb. zinc dimethyl dithic carbamate, said components fluidified in 15 gal. of benzol.

Fat Liquor, Leather

Lecithin	50 lb.
Water	50 lb.
Soda Ash	⅓-1 lb.
Mix the above we	ll and then mix in
suitable quantity of	

Russia Leather from Rejected Hides

The washed and pressed leather is greased in a drum with a mixture of 2 kg. train oil, 5 kg. mineral oil and 4 kg. degras per 62-5 sq. m. of hides, drummed 40 minutes while warm, spread, stoned, dried for 4-5 hours to 38-40% water content and cut through the middle into halves. The damaged spots are cut out, the hides reset and greased by hand out, the inter-see and greated by hand on both sides with a mixture of degras 2 kg., train oil 6 kg., mineral oil 6 kg., lard 6 kg. and tar 5 kg. per 100 sq. m. The leather is left for 12 hours and dried at 28-30° C. to a water content of 32-5%, left for 6 hours to assure a uniform distribution of the water and finally worked over with the whitening sleeker. leather is then dyed, greased on both sides with a mixture of 3 kg. train oil, 4 kg. tar, 6 kg. mineral oil and 2 kg. paraffin, allowed to rest 12 hours, dried at 28-30° C. and treated with a mixture of 150 g. nigrosine, 125 g. gum traga-canth, 50 g. carpenter's glue, 1.5 liter blood and 1 liter milk (all mixed with 12 liters water). The goods are finally dried, polished and sorted.

Protection of Hides and Skins from Skin Beetle

Salt thoroughly applied to hides gives excellent protection against beetle attack. Heavily salted hides which are first rubbed with salt and then soaked in saturated brine for 10 hours or are merely soaked in the brine, are entirely protected during storage for 6 months in the summer in a beetle-infested room. Hides which are rubbed on the flesh side are not so well protected. Hides are protected almost completely by dipping

them, immediately after flaying, in a 2.5% sodium arsenite solution. Spraying sun dried hides on the inside with the sodium arsenite solution does not altogether protect the grain, although it does so to some extent. Sodium arsenite has a marked preservative action on the hides, but a solution stronger than 2.5% is required to prevent decay when hides are dried in the shade in humid regions. When they are stored with salted and untreated hides, the sodium arsenite treated hides do not act as a bait for the beetles and no dead insects are found on them. The sodium arsenite treatment has no deleterious effect on the leather prepared from the hides, and the workmen who handle the hides show no signs of arsenical poisoning.

Stuffing for Welting Leather
Cod Oil 1 gal.
Sulphonated Cod Oil 1 gal.
The above mixture is used per 100 lb.
of welting.

Tanning Shearlings

Soaking: Skins are soaked in clean water, salted skins 10 to 24 hours; dry skins several days, according to condition. Skins must be thoroughly soaked but care must be taken that the wool does not become loose. To prevent this different ingredients are added to the soaks. Small quantities of any of the following may be used: zinc chloride; formaldehyde or alum.

Naphtha or degrading compounds are the most efficient for removing the excess grease; these being reclaimed by distillation and the grease is recovered as a byproduct. In case the stock is not degraded it should be thoroughly washed with a warm soap and soda solution. After degreasing all burrs and brands are worked out. Neglecting to clean out burrs will cause damage in the unhairing machine. Skins are then washed by hand to remove all dirt and to render them as white as possible. This step in the process may be accomplished in the paddle or drum which has a tendency to loosen the wool.

The pickling or tanning may be carried on in the paddle or by hand. If the paddle is used a base solution of approximately 1 lb. of salt for each gallon of water is used and then built up to the desired salometer with equal parts of salt and alum. This amount should be about 4% of each on the weight of stock. This solution may be used several times by the addition of equal parts of salt and alum figured on the weight of the stock.

A small amount of sulphuric acid may be used if desired. This bath is worked up to 50 or 60° C. over a period of 3 days. When stock is struck through it is taken out and drained and is ready for oiling or may be retanned with gambier or quebracho. White and light shade stock is finished out of the alum.

Skins tanned by hand are hest treated on the flesh with salt and sulphuric acid solution. This solution is made with 1 lb. of salt and 1½ oz. of acid to each gallon of water. This solution is applied to the flesh with a brush and the skins piled flesh to flesh or folded down the back with the flesh side in. The next morning the stock is given an alum tan on the flesh made up as follows:

5%
5%
5%
5%
1%
1%

The flour should be worked into a paste, after which the other ingredients are added, the egg yolk being dissolved in a small quantity of cold water. Soda should be added slowly. Two conts of this mixture are given at intervals of 10 to 12 hours at which time stock should be thoroughly tanned. Stock is now thoroughly dried out after which it is sammied back, staked and a light coat of oil given the flesh or a fat liquor may be given, made up of soap, neatsfoot oil and sulphonated oil. Stock before becoming thoroughly dried is staked and stretched. After skins are dried they are restaked, souffed, combed and clipped. If desired stock can then be dyed or blenched.

Russia Leather Odor Bases for this odor are: 2-Tertbutyl 4,5 dimethyl-1-phenol. or 2-Isopropyl-4,5-dimethyl-1-phenol.

LUBRICANTS, OILS, FATS

Gear Lubricant for Arctic Climates

In the northwestern section of the United States and a large section of Canada air temperatures of 40° below sero are not uncommon. At temperatures such as these ordinary winter gear oils are too viscous to permit satisfactory operation of motor cars, and many motor car manufacturers have recommended dluting the gear oil with kerosene to meet these conditions. This practice has always been frowned upon by lubrication engineers since even if the lubricating value of the oil is not entirely destroyed by such dilution, the facilities of the average service station for accurately blending without danger of contamination are not the best. The following formula will produce a lubricant which will give satisfactory performance and adequate lubrication under arctic weather conditions.

Thickened Rape Oil	8 lb.
Asphaltic Black Oil	7 lb.
(90 visc. at 210° F.) Gulf Coast Pale Oil	
Gulf Coast Pale Oil	85 lb.
(100 visc. at 100° F.)	

Sulphur Lubricant Base

The use of sulphur for manufacturing lubricants of high film strength is rapidly gaining popularity. The formula given here will produce a base which can be diluted with mineral oils to make cutting oil and various extreme pressure compounds.

Flower Lard	B u	lpl	ıur		10 90	lb. lb.	
	 _						

Mix well and slowly raise the temperature to 425° F. Maintain mild agitation throughout.

Anti-Rust Compound

Rust and corrosion will do more damage to machinery than several months of hard service. This is particularly true of construction and railway machines which must often be left exposed for long periods. A simple formula for an efficient and economical protective compound is given. The materials should be heated, mixed well and applied with an old paint brush.

Paraffin Wax	6	lb.
Asphaltic Still Residue	94	lb.
(About 1000 visc. at 210°	F.)	

Steering Gear Lubricant

With the general trend to wider treads on automobile tires it has been necessary to redesign steering gear mechanisms to avoid hard steering. Automobile engineers agree that special lubricants are required for most efficient operation.

equitou ros mions omoscus	oporation
Oleic Acid	300 lb.
Lime	43 lb.
Water	16 gal.
Western Cylinder Oil	475 gal.
Sulphur Base	1000 Гь.

Proceed the same as for making lime soap grease except that the sulphur base is not added until the other ingredients are completely cooked.

Mixed Base Grease

The following formula will make a grease which combines the advantages of the smooth texture of calcium soap grease with the cohesive rubber-like character of aluminum cleate. Although the melting point of this grease is not materially higher than a similar calcium soap grease, the melted grease has the slow flowing characteristics of aluminum greases. The formula given is for a medium consistency but other grades can be made by varying the soap content.

Lime	17 lb.
Fat	113 lb.
Aluminum Oleate (Pulp Stock) Pale Oil (100 Viscosity) Water	50 lb. 112 gal. 6 gal.

Place the fat in a steam jacketed kettle equipped with paddles for stirring, add a small portion of the mineral oil, mix the lime with sufficient water to form a thin paste and add this to the material in the kettle. Turn on the steam and start the paddles. When the soap has cooked for 5 hours it should be tested to determine if saponification is completed, if so the steam is turned off and half of the balance of the mineral oil is run into a separate kettle and the aluminum oleate melted in it and this mixture is pumped into the first kettle while still warm. Stirring should be continued until a smooth uniform grease is produced.

Non-Bleeding Grease

One of the difficulties encountered in One of the difficulties encountered in the use of pressure grease is the tendency of the light oil to separate and bleed away leaving the bearing choked with a hard soap. This formula produces a grease which will stand indefinitely without separating. This is not a high melting point grease and is intended for automobile chassis lubrication and similar appliestions. ilar applications.

Green Petrolatum	250	lb.
Paraffin Pale Oil (28° Bé.)		gal.
Lime	9	lb.
Fat	55	lb.
Water	3	gal.

Melt the petrolatum in the mineral oil. Mix well, then proceed as for ordinary calcium soap grease.

Lubricant for Bearings with High Temperatures and Pressure

Formula No. 1

3 g.

Rosin Wool Fat Stearin

Mineral Oil (0.900-7)	80 g.
Castile Soap	15 g.
Caustic Soda (40° Bé.)	4 g.
No. 2	
Rosin	5.5 g.
Wool Fat, Crude	6 g.
Wool Fat, Stearin	11 g.
Tallow	5 g.
Linseed Oil	5 g.
Countie Scale (25° DA)	5 0

Caustic Soda (35° Bé.) Mineral Oil (0.885-90) 78 No. 1 is a high melting fat (150-200° C.), No. 2 melts at about 100° C.

The saponification is done in a directly The saponincation is done in a directly heated kettle (cast iron), which has a removable stirrer, at 150-200° C. Test: should not sweat oil or alkali when pressed with the finger after cooling. If desired, short-out fibers may be added to the mass. Solidify in patterns and cut into briquets.

Metal Rolling Lubricant

micen mountag manison		
Tallow	60	lb.
Yellow Soap	15	lb.
Water	92	gal.
Heat and stir until smooth.		_

Non-Greasy Lubricant

τ	J.	8.	Patent	1,970,902		
Sodium Water	A	lgi	nate	19 100	0Z. 0Z.	
	-	-6.				

Mix to a smooth paste while heating to 100° C. Add Glycerin Boil off nearly all of the water.

Olive Oil Motor Lubricant

Olive Oil (Low Titre)	25	fl.	OZ.
Mineral Oil	75	Ħ.	OZ.

Lubricating Grease for	Carriages
Blue Oil	45 g.
Slaked Lime	6 g.
Rosin Oil	22.5 g. 0.2 g.
Fat Soluble Black Dye	0.2°g.
Dissolve in the blue oil.	_

Chain Lubricant

Formula No. 1

Stearin	85 g.
Beeswax	5 g.
Carnauba Wax	5 g.
Japan Wax	5 g.
No. 2	-
Stearin	85 g.
Beeswax	85 g. 2.5 g.
Carnauba Wax	5 g.
Japan Wax	5 g.
Pour at lowest possible	temperature

and allow to cool slowly and undisturbed.

Penetrating Oil British Patent 414,847

	for	loosening	rusted	metal
parts.				
Engine			1	qt.
Naphth	a or	Keroseno	3	qt.
Carbon	Disu	lphide	2	OZ.
Oil of	Campl	hor	1-2	oz.
Graphi	te, Po	wder	1-4	OE.

O--- 0:1

Core on		
Formula No. 1		
Linseed Oil	300	OZ.
American Gas Oil	600	OZ.
Dark Whale Train Oil	100	oz.
No. 2		
Rosin	200	oz.
Train Oil	200	OZ.
Vulcan Oil	600	oz.

Stuffing Grease

(Melting Point over 96° C.)

Tallow	12	g.
a. Tallow Lard Oil	3	g.
b. Lime Hydrate	2.5	g.
c Zine Orida	2.5	g.

d,	Machine Oil, Refined 4-		
	E. Viscosity 50° C.	79	g.
e.	Yellow Aniline Dyc, Oil		-
	Soluble	0.03	g.
f.	Water	1	
No	otes: Lime Hydrate-M	lade u	p of
fines	t commercial lime hydra	te, dil	uting

with water 1:4. Work bringing a into kettle with 1/3 to 1/3 of needed a; heat to 80-90° C., add slowly b, continue warming. At 100° C. the mass starts "rising" in the kettle, then diminishes when water is evapo-

rating.

Tests: Should be resistant against not too strong finger-pressure; weakly brittle, should not sweat out water or oil when cooled. On the other hand, a water insufficiency is indicated if mass is too brittle (in this case add little boiling water). If tests are satisfactory, add the remainder of d, at 70° C, or warmer—not too slowly, not too quickly. The aniline dye dissolve in mineral oil.

Let stand over night. Stir till cool next day.

Cutting Oil

Formula No. 1 a. Mineral Oil (Spindle Oil) 80 "Tall Oil," Refined 20

"Tall Oil," Refined 20 g.
b. Caustic Potash (40° Bé.) 6 g.
c. Methylhexalin 1-2 g.
Saponify a with b, clear with o.

No. 2

110. 2	
Paraffin Oil (28 to 30° B6.)	250 g.
Rosin	22 g.
Oleic Acid	22 g.
Caustic Soda	3 g.
Water	10 g.
Alcohol	7 g.
No. 3	J

Lard Oil (No. 1) 1 gal. Paraffin Oil (28° B6.) 52 gal. Manipulation: Mix at room temperature.

No. 4

Lard Oil (No. 1) 5 gal.
Extra Lard Oil 5 gal.
Paraffin Oil (28° Bé.) 42 gal.
Manipulation: Mix at room temperature.

Non-Corrosive Cutting Oil U. S. Patent 1.979.250

U. S. Patent	1,979,250	
Mineral Oil	71-74	lb.
Castor Oil	814-914	lb.
Rapesecd Oil	814-914	lb.
Caustic Potash	814-914	lb.
Soda Ash	0.6-11/	lb.
Mix and dilute with	water.	

Brake Oil (Non-Rancid)

a. Mineral Oil (Spindle Oil) 1000 g.
b. Paratoluol Sulphochloride 5-6 g.

or
a. Rape Seed Oil 900 g.
(Camphor Oil 100 g.
b. Paratoluol Sulphochloride 5-6 g.

Dissolve b in little part of a, then add to the above amount.

Gasoline Motor Lubricant British Patent 423,441

Mineral Oil 99 lb. Chromium Oleate 1 lb.

Radiator Anti-Rust Compound

In the past year the automotive industry has given much attention to the prevention of rust and corrosion in automobile cooling systems. Engines with aluminum composition cylinder heads have received the most attention but even in the case of ordinary steel parts it has been found that cooling systems are more efficient if rust and scale formation is prevented.

For this purpose soluble cutting oil such as is used for machining metal is very efficient. The only limiting factors are acidity and alkalinity. Soluble oils having a high acidity will corrode the radiator while too much free alkali will damage aluminum cylinder heads. Several of the formulæ given in volumes one and two of THE CHEMICAL FORM-ULARY will be very satisfactory as cooling system corrosion preventatives. The usual quantity used is ½ oz. of soluble oil for each gallon of water.

Greaseless Lubricating Pencil

Useful for lubricating hinges of automobile doors, etc., as it will not run off and produce stains or accumulate dust.

	mooding and a differ
Beeswax	80 g.
Diglycol Stearate	20 g.
Graphite Powder	100-200 g.
Melt together and	stir until just cold
enough to pour. Po	our into molds and
allow to set.	and moral and

Dynamo Brush Lubricant

Ceresin	20 g.
Tallow, Acid Free	10 g.
Wool Fat, Neutral	10 g.
Castor Oil	10 g.
Vaseline Oil	50 g.

Melt together and add enough organic solvent (Heavy Benzoline, Naphtha or Tetralin).

Soot

5 OE.

Cotton Spindle Machine	Oil
Spindle Oil, Refined (5-6° E at 20° C.) Rape Seed Oil	85 gal. 15 gal.

Veneer Press Caul Lubricant German Patent 596,345

Neutral Soap Lanolin Petrolatum, Liquid Formaldehyde	35 30	0Z. 0Z. 0Z.
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Transformer Oil U. S. Patent 1,988,299

Crude Mineral Oil Phenyl Alpha Naphthylamine 0.5 g.

Transformer Oil Canadian Patent 353,332

To a mineral oil of iodine value of 7 to 20 about 0.5% phenyl a naphthylumine is added to retard sludge formation.

Petroleum Proof Valve Lubricant Citric Acid, Anhydrous Tetraethylene Glycol

Heat at 180-185° C. for 90 minutes; cool. Do not overheat or an infusible moduct will form.

Rubber Mold Lubricant

Cocoa soapstock, a material containing a large percentage of coconut oil saponi fied with alkalies to give a pure hard soap, makes a suitable product for lubricating molds to prevent sticking of the vulcanized stock. If properly made, with out traces of sodium silicate, it will not cause caking on the molds. The recommended quantity is 8 to 12 lb. to a 55 gal. drum of water. The soap is dissolved in water by cooking, either by open steam or external heat of some kind. For easy spraying the solution is kept warm by steam or a small electric heating unit can be applied at the spray nozzle to prevent clogging.

Screw Thread Lubricant

Flaked graphite mixed with a medium grade of lubricating oil to form a paste and applied to the threads of screws and bolts facilitates the backing off of nuts and the removal of screws and machine bolts. The paste, which also is suitable for pipe joints, prevents rust.

	Vacuum	Tap	Grease		
Rubber				30	OK.
Rosin				15	0 %.
Pine Pi	tch			50	OZ.

Journal Grenan

U. S. Patent 1,989,196

5.9	lb.
40.5	lb.
13.1	lb.
6.1	lb.
	5.9 34.4 40.5 13.1 6.1

Spring-Leaf Lubricant British Putent 414,948

White Lend in Linseed	Oil	
(92 Lead, 8 Oil)	83-84	lb.
Graphite Powder	5.2	lb.
Petroleum Grease	10.4-10.5	lb.
Glycerin	0 - 1.3	lb.

Nickel and Monel Drawing Lubricant A paste made of castor oil and lead, recommended for use as a lubricant in the cold forming of Monel metal and nickel, can be removed by a number of solvents. Carbon tetrachloride, being non-inflammable, is to be preferred. Benzene, gasoline, and alcohol produce satisfactory results.

Cold soap and caustic solutions are

not entirely satisfactory but can be used as an alternative, if necessary, when they are used hot.

Wire Drawing Lubricant U. S. Patent 1.944.273

Sodium Alginate	1	lb.
Tallow	4	lb.
Soap	2	lb.
	95	lb.

Drawing Die Lubricant for Diamond Dies

Rye Flour	6	lb.
Water	100	lb.
Beef Tallow	21/2	lb.
Soft Soap	21/2	lb.

Heat and stir until uniform.

Corrosion Protecting Grease

Neutral Petroleum Grease 100 07. Zinc Chromate Powder 21/2 oz. Pyridin Bases (Crude) 1 Rub together to form smooth grease.

Lubricant for Preventing Corrosion French Patent 778,792

Sodium Peroxide	1/4	oz.
Methanol	2	oz.
Hydrogenated Phenol	4	0Z.
Lubricating Oil	100	oz.

Lubricating Haulage Ropes

Before the lubricant is applied, the surface of the rope should be cleaned and dried, because oil or gresse applied to the surface of a rope covered with mud or coul dust, water and old oil will be thrown off without having the slightest chance of penetrating to the interior. In most cases the treatment can be given to the tope during an alle shift.

Main ropes used on inclines can be treated as follows: The rope should be wound very slowly on to the drum, the surface being cleaned as it enters the engine house. Cleaning should be done with wire brushes without using a solvent, such as petrol or paraffin. The brushes may from time to time be washed in paraffin, but this should be shaken off before using the brush on the rope again. The cleaning may be completed with waste or sacking. No solvent (petrol or paraffin) should be used on the rope, because experience has shown that the solvent readily penetrates into the middle of the rope and rapidly dissolves out any remaining lubricant. The rope should be allowed to remain on the drum long enough to allow it to dry as much as possible.

When the rope has been cleaned and dried, the lubricant should be applied by hand with a fairly stiff brush. Devices in which the rope is caused to pass under a roller in a bath of oil are less effective and are wasteful. It is important that the rope should be dry when the lubricant is applied otherwise the oil will not adhere, and the work should be done within the engine house as the rope leaves the drum. If the lubricant is applied in the open, a shower of rain may render useless the whole operation of cleaning and drying the rope. The successful lubrication of a hanlage rope calls for a good deal of skill and patience, but unless it is properly done the time and materials are wasted. It is better to do a portion of the rope well each week than to waste a lot of grease by applying it to the whole of the rope without cleaning and drying.

It is not possible to lay down any fixed periods for the lubrication of haulage ropes, because the periods will vary with the working conditions. A rope

which makes a large number of journeys on a wet incline will need lubrication every week, whereas a rope which makes only a few journeys in the dry may be kept in good condition by less frequent treatment. Excellent results have been obtained on endless rope haulages where the rope is lubricated continuously. In one instance a light mineral oil is allowed to drip on to the moving rope at the rate of one drop per yard; this rope works on a comparatively clean and dry roadway.

Research is in progress as to the best type of oil for applying to ropes in service. At the moment it would seem that the best results are obtained with a medium heavy mineral oil. The oil must be free from acidity, and should contain no filler or soapy material.

Hot Neck Grease

Asphaltic Residue 10 lb.
Caudle Tar Pitch 20 lb.
Paraffin Cylinder Stock (700 Fire Test) 70 lb.

Heat to 550° F. and blow with air until melting point of 200° F. is obtained. Above is cast into blocks and used for the lubrication of roller necks in steel

mills.

High Temperature Lubricants British Patent 431,066

Lubricants for use at high temperatures, e.g., in internal-combustion engines, consist of lubricating oil in which is dissolved or dispersed chromium or an organic compound thereof, and one or more other substances preventing sludging, e.g., organic compounds of tin and/or lead. Up to 1% of each addition is suitable. For example, 0.5 lb. of chromium oleate, 0.1 lb. of tin oleate, and 0.1 lb. of tetraethyl lead are added to 100 lb. of a compounded vegetable and mineral lubricating oil; or 0.4 lb. chromium oleate and 0.1 lb. of tin oleate to 100 lb. or a paraffinic mineral oil.

Non-Chilling Lubricants Formula No. 1

Mix
Castor Oil 3 cc.
Paraffin, Chlorinated (30%
Chlorine) 7 cc.
Spindle Oil, Russian 190 cc.
This gives a highly cold-resistant, clear

No. 2		Turpentino	8.7 lb.
Mix		Ammonia (28%)	4.4 lb.
Castor Oil	10 cc.	Graplute Powder	30 lb.
Paraffin, Chlorinated (30%			
Chlorine)	10 cc.	Watered 11 Off	
(Heat to 200° C.)		Watersoluble Oil	
Spindle Oil, Russian	80 cc.	Naphthenesulphonic Acids	15 g.
· ·	ov	Olein (or Liquid Wool	
No. 3		Fatty Acid)	5-7 g.
Spindle Oil, Russian	40 cc.	Spindle Oil, Refined (60°	C.) 75 g.
Paraffin, Chlorinated (40%		Caustic Potash (25° Ré.)	intil neutral
Chlorine)	40 cc.	Hexalin and Tetralin (1:1) 3-4 g.
Castor Oil	20 cc.		
		Mineral Oil Soluble Cas	ton Oil
Rod Lubricant			
	0.5	To obtain castor oil wh	
a. Ceresin, Yellow	25 g.	soluble in mineral oil, heat	
Sperm Oil	25 g.	the former with 30 parts of	t tricmore-
Tallow	50 g.	ethylene for 2 hours in a clos	
Melt together.		130° C. The pressure will in	
or		atmospheres. After distilling	on excess
b. Ceresin, Yellow	1 g.	solvent, the resulting castor	OII WIII DO
Spindle Oil, Refined	3-8 g.	soluble in mineral oil. This not be brought about by hea	
	, o 9.	alone or by refluxing with	
Melt at low temperature.		second method is to heat in a	
~		90 parts of castor oil with	
Solid Lubricant		carbon tetrachloride for 2 ho	
Formula No. 1		The pressure increases to	
Canadian Patent 344,	966	atmospheres. Dissolve in min	
Heavy Distilled Naphthenic		distil off excess solvent, remov	
Petroleum	30.8 lb.	traces by distillation in vacuo	
Residual Naphthenic	00.0 10.	,	
Petroleum	13.6 lb.		
Stearic Acid	14 lb.	Lubricant Insoluble in Organ	nic Solvents
Oleostearin	28 lb.	Mix to a paste the following	g:
Caustic Soda	6.6 lb.	Anhydrous Glycerin	25 oz.
Water	7 lb.	Dextrin	7 oz.
No. 2		Pure d-Mannitol	3.5 oz.
Canadian Patent 344,	0.67	Heat carefully with const	ant stirring
•	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	until the solid material is di	swalved and
Viscous Naphthenic		the solution begins to boil, t	hen cool to
Petroleum	43 lb.	room temperature with stirri	
Animal Fat	39.4 lb.	crease the viscosity, add mo	
Aluminum Stearate	4.7 lb. 5.3 lb.	to increase fluidity add more	
Caustic Soda		increase greasiness add more	
Slaked Lime	0.6 lb.	9	
Water	7 lb.		
✓ Hard Grease		Tempering Fats (Bath to (Quench and
	10 -	Harden Steels)	-
Train Oil Fatty Acid	12 g.	Formula No. 1	
Lime, Hydrated	2 g.		E00
Zinc Oxide	2 g.	Peruvian Bark Powder	500 g.
Spindle Oil	82 g.	Neatsfoot Meal	500 g.
Water	2 g.	Halt Sultrator	850 g.
Melting point 75° C.		Saltpeter	250 g. 15 g.
		Potassium Ferrocyanide	1000 g.
Graphite Lubricant		Soft Soap	2000 B.
U. S. Patent 2,003,56	34	No. 2	10 ~
	_	Beef Tallow	10 g.
Degras (Free from Fatty	00 Ib	Potassium Ferrocyanide,	2 ~
Acids)	20 lb. 16 lb.	Powder Wax	2 g. 2 g.
Kerosene Water	16 lb. 75 lb.	Colophony (Rosin)	2 g.
AA Breil	10.	Company (mosm)	~ ₽.

Waterproofing, Perilla Oil

One method is to react one part of straight phenolic resin with 2 or 3 parts of perilla oil at between 500° and 550° F. If polymerized perilla oil is used, even Another better results are obtained. method is to employ some wood oil. For instance, one part of straight phenolic resin to 2 parts of wood oil may be reacted together and then extended with various amounts of polymerized perilla oil. Another formula is phenolic resin 5 parts, wood oil 10 parts, perilla oil 85 Another is 10 phenolic resin, 20 wood oil, and 70 perilla oil. All parts are by weight.

Coloring Lubricating Oils British Patent 424,205

Lubricating oils are improved in color by adding a solution in mineral oil or other blending agent of the product obtained by heating together until fluorescence develops, an acridine, rhodamine, eosine, or eurhodine dye with stearic acid and a water-insoluble soap. Soaps specifled are aluminium stearate, magnesium stearate, oleate, or resinate, and zinc soaps. For example, 1 lb. of phosphine 5G., 1 lb. of stearic acid, and 3 lb. of aluminium stearate are heated to 120° C. until the fluorescence is a maximum; the mixture is cooled, pulverized, and dissolved to a 10% solution in a mineral oil miscible with lubricating oil. 0.25-0.5 gal. of the solution is added to 100 gal. of lubricating oil.

Refining Lubricating Oil U. S. Patent 2,020,954

Stock of about 68 viscosity index is subjected to the simultaneous action of 10% of aluminum chloride and 10% of fuller's earth at a temperature of about 350° F. for 1/2 hour.

Purification of Lubricating Oil

If lubricating oil is shaken with phenol, the lower layer consists of oil and impurities in phenol; the upper layer consists of phenol dissolved in pure oil. The phenol is removed and recovered by distillation or by washing with sulphuric acid.

Dewaxing Mineral Lubricating Oil U. S. Patent 2,014,629

Amorphous wax is eliminated by treating the wax bearing oil with 3 to 10% of substantially anhydrous aluminum

chloride at a temperature of about 200° for a half to four hours, thinning with a light distillate, chilling and filtering.

Dewaxing Oil U. S. Patent 1,978,010

A process for treating wax-oil mixtures comprises mixing 1 to 4 volumes of methylene chloride with 1 volume of the mixture, chilling the mixture to a temperature below 0° F. and filtering precipitated wax from the mixture.

Preventing Discoloration of Oils and

British Patent 410,834

Discoloration of animal or vegetable oils or fats on exposure to air and light is prevented by incorporating not more than 0.05% of colloidal copper, cobalt, cadmium or silver, or of the carbonate of cobalt, copper, lithium, manganese, cadmium, barium, bismuth, the nitrate of calcium, beryllium, or lithium, the acetate of sodium, copper, manganese, the hydroxide or cobalt, beryllium, copper, thorium or of a mixture of cobalt carbonate and copper carbonate with or without bismuth subcarbonate.

Reclaiming Used Lubricating Oil U. S. Patent 1,936,901

Used Lubricating Oil

ned On	1	gaı.
Calcium Hypochlorite	6-8	gal.
Sulphuric Acid	6	lb.
Mix together and then	add:	
Sodium Silicate	50-100	
Water	10-20	gal.
TT 4 -4 FO 1000 CL		_

100 gal.

Heat at 52-122° C. for two hours. Cool; add water, 3 gal., and separate clear oil.

Fat and Oil Bleaching

In refining fats and oils the color is improved by adding 8 to 10% soap stock to the fat.

Decolorizing Tea Seed Oil

Kaolin Animal Charcoal	25 lb. 20 lb.	
The above mixture has		0

Increasing Viscosity of Mineral Oils British Patent 416,513

Thickened mineral oils which form gels at room temperatures are obtained by dissolving less than 2% of cellulose stearate or palmitate in the heated oil.

Oil Filter Mass U. S. Patent 1,940,317

Cotton Waste Curled Hair 75 oz. 25 oz.

Fat Hydrogenation Catalyst
The catalyst is prepared as follows:
Precipitate a solution of 160-300 g. per
liter nickel sulphate with a 15° Bé.

sodium carbonate solution at not ower 32-65° C., filter on a filter press, wash till free from sulphates with water at 30-50°, dry 4 to 5 hours at 100-105°, grind, sieve, mix with sunflower seed oil and reduce by heating the oil in presence of hydrogen; time of reduction is 5 hours; the temperature is raised to 170-200° during the first hour, to 200-240° during the next two hours and to 240-245° during the last 2 hours. Reduction of the catalyst can be carried out in the same autoclave as the subsequent hydrogenation. The activity of the catalyst lasts over a prolonged period.

MATERIALS OF CONSTRUCTION

Metal Cleaning

Many "mysterious" finishing troubles are due to improper cleaning. What cleaning materials and methods to select will depend upon: (1) the size and character of articles to be cleaned, (2) their surface condition, (3) the volume of work to be handled, (4) the kind of finish to be applied, and (5) various conditions peculiar to the particular factory department wherein the cleaning is to be performed.

Rust, dust, greases, and grit can be cleaned off metal surfaces by the use of one or more of several methods. They may be burned off, chemically removed with an acid or an alkali solution, absorbed by gasoline or naphtha, buffed, or removed by sandblasting.

Old varnish or paint may be removed by the burn-off process, preparatory to refinishing. A temperature of 650° to 700° F, is required to dislodge the old coating which can then be wiped off with a rag while still hot. The burn-off (oven) process is also a means of drying washed and chemically treated parts. Heavy rust spots are usually removed

Heavy rust spots are usually removed by wirebrushing, sandpapering or sandblasting. Thin coatings of rust may be removed either by kerosene or gasoline or by pickling in a solution made of commercial sulphuric acid diluted in water. Other solutions used are: (1) A 20% solution of sodium citrate and water, (2) a 10% solution of ferrous sulphate and water, and (3) a 3½% solution of boric acid and water.

Aluminum parts are prepared for a baked finish by a thorough cleaning with gasoline or naphtha, and a subsequent oven-drying. Old paint and varnish may be removed from aluminum with any standard paint or varnish remover.

Metal Cleaning Composition Canadian Patent 345,172

A compound containing trisodium phosphate and sodium dichromate is used for cleaning tin-coated metal. It inhibits checking or spangling. A satisfactory composition contains trisodium phos-

phate 55 lb., sodium carbonate 40 lb., and sodium dichromate 5 lb.

Cleaning Metal Before Painting

Cleaning	metan	Defore	Lamming
Apply Ammonia Alcohol Water	(28%)		1 l. 26 l. 25 l.

Wipe off metal thoroughly after application.

Cleaning Iron and Steel

U. S. Patent 1,943,875

Prior to galvanizing or tinning the metal is exposed to the fumes of 1 to 2% of phosgene at 100-200° C.

Cleaning Tin Surfaces

a. A bath is made up of palm oil that has been heated to 300° F. Any method of heating may be employed as the flash point of the palm oil is quite high. Generally speaking, there is no danger of overheating. Probably the most practical method of heating is by using a steam coil in the palm oil container, as the temperature may be easily controlled.

The work is dipped into the solution of heated palm oil for two to three minutes and removed. No further processing is required for the palm oil is quite liquid at this temperature and will flow freely from the work. It may be found necessary to remove some of the oil by using an air blast to blow the oil from the work.

The method suggested above will operate well on small work. However, if the work is large, it may be necessary to preheat the work before immersing it into the oil bath. Without preheating heavy work, the oil will cool too quickly when the work is being removed from the solution and will leave an unsatisfactory waxy deposit on the work. The preheating is best accomplished by immersion in superheated water long enough to heat the work sufficiently. Upon removing the work from the heated water, it may be immersed immediately in the palm oil bath.

b. Another method that may be used with good results is to immerse the work in a 2% solution of water and nitric acid. This procedure is most efficient if the work is first preheated in water as suggested in Method a. The acid dip is immediately followed by immersion in a rinse of kerosene oil. The duration of the acid dip must be found by experiment as the length of dip depends upon the thickness of the oxide. This may be easily determined by the trial and error method. Too short a dip does not restore the luster, and too long a dip increases the tarnish and produces a spangle effect as in galvanizing. The acid dip and kerosene rinse are operated at room temperature.

The drying of the work is best accomplished by drying in heated sawdust. Care must be exerted in this operation as machined work will rust if it is not dried

thoroughly and quickly.

The success of cleaning of tinned work depends upon the quality of the tinning that was on the work originally. It is impossible to produce a luster on an article that had a poor finish in the first place.

Cleaning Monel Screw Machine Parts

The use of sulphur base cutting oil in high speed automatic screw machine operations, may discolor the Monel metal parts. This discoloration is due to the formation of metallic sulphides by the

sulphur in the oil.

The discoloration is readily removed by dipping the parts in a cold solution of sodium cyanide. The solution is made up in the proportions of water 1 gal., sodium cyanide 1/2 to 1 lb. The time required for cleaning is from 5 to 30 minutes, depending on the degree of dis-coloration. Caution should be used in handling this solution as it is a deadly poison.

Coloring Metals

Metals are colored chemically or electrochemically by producing thin films of oxide, sulphide, phosphide, silicide, nitride and carbon on their surface. For quantity production, coloring is usually carried on in a rotating drum, while large pieces and objects of art are treated by hand. A few recipes follow:

1. For Copper

a. Brown: immersing in molten sodium nitrate, or imbedding in a paste of 15 parts ammonium carbonate and 5 parts each of copper acetate, tartaric acid in

vinegar, and salt; another solution is 25% copper sulphate, 25% nickel sulphate, 12% potassium chlorate, 7% potassium permanganate.

b. Gray-black: a hot watery solution of 12% copper sulphate and 1% potas-

sium permanganate.
c. Black: 40-50° C. (104-122° F.) warm solution of 600 g, copper nitrate in 200 g, water and 2.5 g, silver nitrate in 10 g, water is brushed on the object and dried at 230° C. (446° F.); or a solution of 10% sodium chlorate, 5% caustic soda and 10% potassium persulphate is used for immersion.

d. Green patina: solution of 25% ammonium chloride, 25% ammonium carbonate, or an acctic acid with an addition

of 1-2% tartaric acid.

e. Blue: 80° C. (176° F.) hot solution of 13% thiosulphate and 3.5% sugar of lead, or of 100 g. potassium chlorate, 100 g. ammonium nitrate and 1 g. copper nitrate in 1 l. water. The objects are immersed for 5-10 minutes,

f. Purple-gray: immersion in a solution of antimony trichloride in water with an addition of equal weight of 5% hydrochloric acid.

2. For Zinc

a. Yellow: aqueous solutions of 5% copper sulphate, 5% sal ammoniac and 3% ammonium chloride are brushed

b. Black: solution of 16% copper sulphate, 8% potassium chlorate in 1 l. water; or a cold solution of 8 parts hydrochloric acid, 3 parts copper chloride, and 2 parts copper nitrate in 64 parts of

water,

c. Iridescent: immersing in a solution of 3 parts tartrate of copper oxide and 4 parts of caustic soda in 48 parts of water. According to duration of immersion, purple, blue, green, yellow or red hues are obtained.

d. Purple: immersing in a warm bath -60° C. (140° F.)—of 60 g. nickel ammonium sulphate, 60 g. ammonium

chloride, 1 l. water.

e. Steel blue: a bath of 60 g. cobalt ammonium sulphate, 60 g. ammonium chloride, 1 l. water.

3. For Tin

Tin, before coloring, is either copperor brass-plated and then treated as given for these metals.

4. For Aluminum

Aluminum can generally be colored black only, either by burning in a layer of carbon produced by linseed oil or albumen, or by immersing in a 5% platinum chloride solution in water or 1% platinum chloride solution in alcohol, and left to dry in 150° C. (302° F.). The methods used for black-coloring of copper can also be applied.

5. For Iron

Black can be obtained by burning in linseed oil, tallow or wax at 400° C. (752° F.) in rotating drums, or in aqueous solution of 2% copper chloride, 2% bismuth chloride, 4% mercury chloride, 12% hydrochloric acid and 10% al-cohol; the object is boiled in this solution. Iron can be burnished at 100° C. (212° F.) in a solution of 1% ferrous chloride, or 7% ferrous chloride and 0-2% mercury chloride with addition of a few drops of hydrochloric acid. A red-dish-brown is obtained by applying a solution of 15 g. ferric chloride in 1 l. water and leaving it in for a few hours.

6. For Silver

Black is obtained by either a 1% aqueous solution of ammonium sulphide or a 5% solution of ferric chloride and rinsing in 2% caustic soda.

7. For Gold

A red-gold tint is produced by a warm solution of 115 parts salt, 230 parts saltpeter, 170 parts hydrochloric acid and 150 parts water; or of 3 parts hydrochloric acid, 1 part nitric acid, 2 parts salt in 40 parts water.

8. For Nickel

Treating with platinum chloride or sal ammoniae containing ammonium sulphide gives black and gray tints.

Black Finishing Chromium Plate U. S. Patent 1,937,629

Immerse articles for	20-30 minutes in:
Sodium Cyanide	45 lb.
Soda Ash	35 lb.
Salt	20 lb.

at temperature of 700-900° C.

Coloring Copper a Green-Blue

A malachite coating is formed on a copper anode in an aqueous solution of an alkali carbonate (8% sodium bicarbonate), using a c.d. of 1-20 amp./sq. dm. The coating may be applied to copper roofs, etc., by means of a cloth-covered roller soaked in the electrolyte. The coating is green and adherent, and changes to brochantite within a year without flaking.

Coloring Brass

Cheap Rose Gold Finish

The work which must be brass is placed in the following dip until a smut is produced:

Copper Sulphate	16	oz.
Muriatic Acid	₩	gal.
Water	1	gal.

Dissolve the copper sulphate in the water and then add the acid. The work should have a deep red smut which should be lightened somewhat by placing in a saturated salt solution for a few seconds. Plate in the regular fine gold solution, then relieve the high lights with bicarbonate of soda, replate in gold solution for a few seconds, dry and lacquer.

Blue Black Color

Copper Carbonate	1 lb.
Ammonium Hydroxide	1 qt.
Water	3 at.

Add the water after the copper carbonate and the ammonia have been thoroughly mixed. Use at a temperature of 175° F. and immerse the work until the color is obtained (usually from ½ to 1 minute). There must be excess copper carbonate.

Verde Finishes

Formula No. 1

White Arsenic	8	0 Z.
Muriatic Acid	1	qt.
Copper Acetate	2	lb.
Copper Carbonate	⅓	lb.
Ammonium Chloride	2	lb.
Water	2	gal.

Dissolve the arsenic in the muriatic acid with the aid of heat and then add the copper carbonate. Dissolve the copper acctate and the ammonium chloride in the water and mix the two solutions thoroughly. This is used with a brush. If desired as an immersion, reduce to twice the volume with water.

No. 2

Copper Acetate	4 oz.
Copper Acetate Copper Nitrate	4 02.
Ammonium Chloride	4 oz.
Water	1 gal.
No. 3	- 0
Copper Nitrate	8 oz.
Ammonium Chloride	4 oz.
Acetic Acid	4 05.
Chromic Acid	1 02.
Water	1 gal.

Apply lightly with brush and let dry. If finish is not even, brush again with the verde solution and let dry.

Verde Color

(1111411) (110011)	
Copper Sulphate	8 oz.
Ammonium Chloride	4 oz.
Sodium Chloride	4 oz.
Zinc Chloride	1 oz.
Acetic Acid	2 oz.
Water	1 gal.
m. 33111 - 4 1 - 4	almorin Wi

The addition of 1 oz. of glycerin will prevent the green from drying too fast and produce a more even color. This solution is used for immersion and if the color is not uniform, repeat immersion as many times as desired, allowing the work to dry thoroughly between immersions.

Electrolytic Verde Finish

Potassium Bichromate	8 oz.
Copper Sulphate	12 oz.
Water	1 gal.

Use solution at a temperature of 80° F.; lead anodes and 8 to 10 volts. Then set color in an alkaline solution.

Brown on Brass Formula No. 1

Golden Sulphuret of	
Antimony	4 oz.
Caustic Soda	8 oz.
Water	1 gal.

Use as near the boiling point as pos-

Scratch brush dry. If the color is not dark enough, pass through a dip composed of 2 oz. sulphuric acid, water 1 gal.

No. 2	
"Liquid" Sulphur	1 oz. 1 gal.

The work is immersed in this solution for a minute or so and then without rinsing immersed into a solution made of sulphuric acid 1 oz., nitrie acid 1 oz., water 1 gal. If color is not dark enough, repeat both dipping operations and scratch brush dry.

Blue Color on Brass

Hyposulphite of Soda Lead Acetate Water	4	oz. oz. gal.
11 1000		

Use at boiling temperature and immerse just long enough to produce blue color.

Green Color on Brass

GIOCH COLOI ON DIMES		
Nitrate of Iron		05.
Hyposulphite of Soda		OE.
Water	1	gal.
Use boiling temperature.		

Verde Color on Brass

A CLOS COLOS OR ANGRO	
Copper Nitrate	16 oz.
Ammonium Chloride	4 oz.
Acetic Acid	1 qt.
Water	3 qt.
- 41 1.4	T

Immerse the work and let dry. If color is not uniform use a painter's sash brush which is moistened with the solution and stipple lightly.

Old English Finish on Brass

Two solutions are necessary to produce this finish, one a sulphur solution, the other an acid solution.

Formula No. 1

Liquid Sulphur	⅓ oz. 1 gal.
Water No. 2	ı gan
Copper Sulphate	2 oz. 1 cal.

The work is thoroughly cleaned in an alkaline cleaning solution, then dipped in No. 1 solution, and without rinsing dipped in No. 2 solution. These dips are only momentary. Rinse in clean cold water and repeat dipping operations until a light color is produced.

For an even finish, scratch brush, dry and repeat dipping operations in solutions No. 1 and No. 2; finally scratch brush dry and lacquer.

Coloring Brass or Copper

(Use Brush or Immersion)

Diaca	
Potassium Sulphide	2 oz.
Ammonium Chloride	2 lb.
Water	1 gal.
Brown	•
1310MT	
Ammonium Sulphide	2 oz.
Water	1 gal.
Blue Green (180° F.)	_
Sodium Thiosulphate	1 oz.
Iron Pernitrate	8 oz.
Water	1 gal.
Rust Brown	
Barium Sulphide	2 oz.
Water	1 gal.
Red (120° F.)	_

Red (120° F.)	
Sulphate	4 oz. 2 lb.
	1 gal

Verde Green (75° F.) Copper Nitrate 5 oz. Ammonium Chloride 5 oz. Chloride of Lime 5 oz. Water 1 gal.

Coloring Bronze

Formula No. 1

Use a boiling or near-boiling solution containing 50 to 60 g. copper sulphate per liter of water. Additions of alum (potassium aluminum sulphate) give colors tending toward the violet red. About 20 g./l. are recommended.

Additions of verdigris give olive-green colors. About 30 g./l. are recommended, with further additions of 5 to 10 g./l. if desired.

A very pretty red may be obtained from the following:

Copper Sulphate	62.5 g.
Verdigris	10 g.
Alum	25 g.
Water	ĩ ĩ
Acetic Acid	few drops
Exact reproduction	 thin a-lam :

reproduction of this color is sometimes difficult.

No. 2

Bronze may be colored in the following:

Sodium Chlorate	50 g.
Copper Sulphate	125 g.
Water	1 l.

If copper nitrate is used instead of copper sulphate, less sludge is obtained. 148 g. of copper nitrate should be used. The following colors are obtained:

Solution near boiling-greenish goldbrown obtained in 5 minutes.

Solution near boiling-gold brown obtained in 10 minutes. Solution cold-yellow brown obtained

overnight. The effects of additions are as fol-

lows: Addition of ferrous sulphate-slight

change toward olive green.

Addition of ferric ammonium sulphate similar to above but lighter in color.

Addition of ferric sulphate similar to

above but with strong etching.
Addition of nickel sulphate—increase in yellow brown.

Addition of ammonium sulphate-lighter color and more yellowish brown, partly toward greenish.

No. 3

Antique Green-Oxidized Effect

After cleaning, dip and/or brush with stippling effect, using the following so-

mon:	
Water	1 gal.
Iron Chloride	3 oz.
Sal Ammoniac	16 oz.
Verdigris Powder	8 oz.
Common Salt	10 oz.
Cream of Tartar	4 oz.

No. 4

If bronze is being exposed to the atmosphere, rub it with cotton waste soaked in boiled linseed oil to obtain, on aging, a dark brown adherent color.

No. 5

For brown, reddish bronze, or blueblack tones use:

Water	1 gal.
Liver of Sulphur	2 oz.
Caustic Soda	3 oz.
Use a temperature of 160°	to 180° F

Coloring of Copper

termines the color.

The pieces to be colored are first cleaned of all oil and grease with gasoline and then lightly etched in the following solution:

Water Concentrated Sulphuric Acid 10 oz.

They are then thoroughly washed in water before immersion in one of the following coloring solutions.

Brown to Steel Blue Color

Liver Salt	of	Sulphur			g.
Water				100	g.
Thie	hath	works	hattar	mhan.	_

This bath works better when kept warm. The pieces are left in the bath until the desired color has been obtained.

Gray-Brown Color

Iron Chloride	3	g.
Water	100	
The pieces are heated and dipp	ed.	_

Brown Colon

DIOMT COIOL	
Powdered Copper Sulphate	100 g.
Zinc Chloride	100 g.
Water	200 g.

This forms a paste which is smeared over the surfaces to be colored and allowed to dry.

	MATERIALS OF
Other Brown Colo	ring Solutions
Liver of Sulphur Carbonate of Ammo Water	5 g. nia 10 g. 250 g.
Copper Acetate Ammonium Chloride Ammonia (10%) Vinegar This is brushed on.	10 g. 5 g. 25 g. 160 g.
Old copper effects brushing sulphuric ac sions and thoroughly the desired amount of been formed. After the colored thoroughly washed and	id in the depres- washing off after f green oxide has pieces have been dried they should
be polished and given	i preservative coat

en ld b at of a suitable lacquer or the following mixture:

Carnauba Wax	100	g.
Japan Wax	100	g.
French Turpentine	1000	g.

Coloring Copper Formula No

rorman no. 1	
Potassium Chlorate	1 oz.
Copper Sulphate	4 oz.
Water	1 gal.
Use hot, scratch brush wet.	If color

is uneven, repeat coloring operation and scratch brush dry.

No. 2

A darker or more red color is produced in this solution.

Copper Sulphate			4 oz.
Nickel Sulphate			2 oz.
Potassium Chlorate			1 oz.
Water			1 gal.
Finishing operations	are	the	same a

above. No. 3

Various shades of bronze from a chocolate color to a black can be produced in this solution.

Potassium Water	Sulphide	1/2	to 1	oz. gal.

For the light shades use cold and a short time of immersion. For darker, use hot, with longer immersion.

No. 4

Various colors are produced in any of the following solutions used either hot or

Yellow Barium Sulphide Water	1 oz. 1 gal.
No. 5	ŭ
Yellow Barium Sulphide	1 oz.
Calcium Sulphide	1/2 fl. oz.
Water	1 gal.

No. 6				
Golden Sulphuret of				
Antimony	1/2	to	1	OZ,
Caustic Soda	1	to	2	OZ.
Water	4		1	gal.
No. 7	¥			
Copper Sulphate		:	12	oz.
Acetic Acid			4	OZ.
Caustie Soda			4	OZ.
Water			1	gal.
No. 8				
Copper Sulphate			4	OZ.
Copper Acetate			2	OZ.
Potassium Chloride			6	oz.
Water			1	gal.
No. 9				
Copper Sulphate			8	oz.
Potassium Permanganat	o		1	OZ.
Water			1	gal.
-	-			

Coloring Silver

Formula No. 1 Sulphide Coloring

Dip in solutions of sodium or potassium sulphide.

No. 2

Tellurium Black

Dissolve 1 oz. of pure tellurium dioxide in 16 oz. concentrated hydrochloric acid to which have been added 8 oz. water. Boiling the solution will probably be necessary.

The solution so obtained should be diluted with water, the amount depending on the anticipated use. For brushing, use about 1 part of the above with 2 parts water. For dipping, a much weaker solution is advisable.

Better results are obtained from a hot than from a cold solution.

No. 3

Platinum Black

Silver placed in hot 5% platinic chloride solution rapidly turns jet black.

No. 4

Iron Oxide Finish on Silver

Immerse the silver for about 5 seconds in a solution containing 1200 g. ferric chloride per l. water. Rinse the article and immerse for 15

seconds in a solution containing 20 g. caustic soda per l. water.

Better results are obtained if the ar-

ticle is made the cathode in the latter solution.

No. 5

Black Nickel

For relief designs on silver, black nickel is often used. The presence of

zinc or copper in a nickel plating solution will cause distinct darkening of the nickel deposit. A simple formula is: Water 11

Water
Nickel Ammonium Sulphate
Ammonium Thiocyanate
Zinc Sulphate

1 1.
50 g.
10 g.
6 g.

Carbon anodes are used, and the silver article is made the cathode at about 3 amperes per sq. ft. Excess black nickel is removed with a tampico wheel and pumice.

No. 6

Pink Color on Silver

A pink color may be given silver by immersing it in a hot solution of copper chloride,

Antique Silver Finish Formula No. 1

Roughen surface (as by acid dipping) and then dip into the following solution:

Lead Acetate 3 g.
Sodium Thiosulphate 140 g.
Water 1 1.

Temperature 140° F.

No. 2

Dip article into following solution:

Ortho Arsenic Acid	50 g.
Sodium Carbonate	20 g.
Potassium Cyanide	25 g.
Water	1 Ĭ.

Add the chemicals to the water in the above order, with thorough mixing of each.

No. 3

Dip article into solution containing 15 g. potassium sulphide per l. of water. Rinse in water and dip into following:

Copper Sulphate 9 g. Sulphuric Acid (Conc.) 3 g. Water 1 i.

Polish article with fine pumice and dip into weak solution of potassium cyanide containing sodium hydroxide.

Imitation Antique Silver Finish

An imitation antique silver appearance may be given iron, for example, by first cadmium plating it, and then dipping it in the following:

um Chlorate Nitrate	·	60 g. 40 g. 1 l.

Preventing Flaking in Steel Flakes, especially in steels of the S.A.E. 3312 type, can be avoided by thoroughly deoxidizing before adding the iron alloys, by mixing the bath well, by pouring at 1420-50°, by slow cooling and heating in the range 300-700°, and by forging at high temperature.

Coating Iron with Aluminum British Patent 432.212

Iron wire is exposed to ammonium chloride vapors at 500-700° C. and passed directly into a bath of molten aluminum.

Phosphate Coating for Steel Canadian Patent 351,060

Sodium Nitrate 100 lb. Manganese Acid Phosphate 115 lb. Copper Carbonate 19 g. Water 400 gnl.

Coating Steel with Zinc Phosphate U. S. Patent 1,926,265

Dip steel in:

Zinc Cyanide 3 lb.
Zinc Acid Phosphate 15 lb.
Water 100 lb.

while heated at 75° C.

Foundry Parting Powder British Patent 412,931

Kieselguhr 92-97.5 lb., wax 6-2 lb. and resin 2-0.5 lb, the kieselguhr being thoroughly mixed with the molten wax and, after cooling, the mixture being ground with the powdered resin.

Improving Malleable Iron Castings U. S. Patent 2,024,014

The process for the heat treatment of malleable iron castings containing 0.6 to 5% copper comprises heating the malleabilized castings to a temperature in the range of approximately 700 to 850° C.; cooling at a rate greater than approximately 25° C. per hour to a temperature in the range of approximately 400 to 600° C.; and without further cooling maintaining in that temperature range for sufficient time to produce a substantial increase in hardness.

Increasing Carbon Content of Iron U. S. Patent 2,021,159

Add to molten metal after leaving cupola a mixture of:

Sodium Nitrate 20 lb. Carbonaceous Material 80 lb.

MATERIALS OF	CONSTRUCTION	223
Case Hardening Composition Formula No. 1 U. S. Patent 2,002,180 Sodium Cyanide 9 lb. Barium Chloride 6 lb. Barium Carbonate 8 lh. Calcium Fluoride 2 lb. No. 2 U. S. Patent 1,952,090 Calcium Chloride 20 lb. Salt 10 lb. Sodium Cyanide 0.15-0.3 lb.	Salt Soda Ash Ammonium Chloride Barium Carbonate Potassium Dichromate No. 3 British Patent 416,1 Coat with following and h burizing temperature: Carbon Powder Barium Carbonate Nickel Steel (20%) Turnings Ashestos Fiber Sodium Siliente (d. 1.33)	
U. S. Patent 1,942,937	No. 4	
Heat metal at 1010-1065° C. in a mix-	Patented	
ture of:	Immerse in a fused sult ba	
Charcoal Powder 40 lb. Hardwood Sawdust 24 lb. Manganese 20 lb. Chromium 5 lb. Borax 8 lb.	Bodium Mitrate Barium Carbonate Salt	15-40 lb. 20-40 lb. 10-15 lb. 5-10 lb.
Chopped Pea Plants 3 lb.	Temperature is maintaine 960° C, and a current of amn	nonia gas is
allowing free access of air.	passed through the bath to nitride case.	produce a
No. 4	A * . TT 1	,
British Patent 412,173	Air Hardening Stee U. S. Patent 1,976,3	
Metal is dipped in following:		_
Ground Rice	An air quenched article of is composed of about 3.1 to 0.25% carbon, 1.5 to 2% manganese, the ba substantially all iroz.	4% copper, and about
After drying the coated metal, heat to 900-950° C. in a non-oxidizing at-	Hydrogen Chloride Resists German Patent 596,0	23
mosphere.		-74 kg. -25 kg.
Hardening Steel		-25 kg. -14.5 kg.
Formula No. 1	Tantalum 0.5	-7 kg.
Austrian Patent 142,401		-1.5 kg. -8.5 kg.
Potassium Ferrocyanide 70-80 kg. Soda Ash 2-5 kg.	Molybdenum 0.4	-7 kg.
8alt 6-12 kg.	Silver 0.1	-4.5 kg.
Acetylene Carbon 3-8 kg.		
Potassium Carbonate 2-3 kg. Ammonium Chloride 2-3 kg.	Surface Carbonization o	
Gum Arabic 2-3 kg.	U. S. Patent 1,950,1	
The above mixture is strewn over the steel which is then heated.	Etch surface in 15% nitric dry; heat at 900° C. in a l vapor.	hydrocarbon
No. 2	Anti-Carburizing Comp	osition
U. S. Patent 2,016,477	U. S. Patent 1,982,	
Soybean Powder 90 lb. Sodium Cyanide 3 lb.	Copper Chloride Oxalic Acid	2 lb. 3 lb.

Lead Oxide Copper Sulphate Water	1 lb. 5½ lb. 5 lb.
Metallographic Etchir Copper Ammonium Chlor Hydrochloric Acid Ferric Chloride Water	
Etching Hardened	Steel
Mercuric Nitrate Nitric Acid Water	5 oz. 38.5 oz. 89.5 oz.
Etching Stainless Formula No. 1	
Nitric Acid Hydrochloric Acid Denatured Alcohol Water Solution used cold.	32 oz. 3 oz. 16 oz. 96 oz.
No. 2	
Ferric Chloride Hydrochloric Acid Water This solution may be u 120° F, or electrolytically.	20 g. 20 g. 60 cc. sed warm at
Steel Pickling Inhii U. S. Patent 1,932	
Di-o-tolylthiourea Evaporated Waste Sulphi	4 lb.
Liquor	6 lb.
Salt Soda Ash	10 lb.
BOUR ABR	1 lb.

The above is formed into blocks.

Metal Pickling Inhibitor Canadian Patent 353,320

Pyridine 80 g.
Benzyl Chloride 140 g.
Heat to 160-170° C. and cool to 75100° C. and then dilute with any solvent.

Ore Briquettes for Open Hearth

100 lb. 10 lb. 1 lb.

More satisfactory results are gotten by using above briquettes than when using dust ore.

Age Hardening Silver U. S. Patent 1,984,225

Sterling silver capable of age hardening to a hardness of from 84 Rockwell B to 94 Rockwell B consists of pure silver at least 92.5%, copper 2.5 to 7.4% and aluminum 0.1 to 5%.

at least 92.5%, copper 2.5 to 7.4% and aluminum 0.1 to 5%.

A process of making sterling silver articles of a hardness of from 80 Rockwell B to 94 Rockwell B consists in first alloying at least 92.5% silver, from 7.4 to 2.5% copper and from 0.1 to 5% of a metal selected from the group consisting of aluminum, magnesium, lead, antimony, and beryllium, then fabricating the article to form by known cold working operations, then subjecting the article to a preliminary anneal and quench from about 1150° F. to 1400° F. and finally subjecting the article to an age hardening heat of about 570° F. for about one hour.

PHYSICAL PROPERTIES OF METALS

	_		Melting	Point	Weight
Metal	Specific Gravity	Specific Heat	Deg. Cen- tigrade	Deg. Fah- renheit	in Lbs. per Cubic Inch
Aluminum:					
(Cast)	2.56	.2185	658	1217	.0924
(Rolled)	2.71	• • • •	• • • •	• • • •	.0978
No. 38 Alloy (Rolled)	2.74	• • • •		• • • •	.0989
No. 12 Alloy (Rolled)	2.82	• • • •	624	1156	.1018
Antimony	6.71	.051	630	1166	.2424
Bismuth	9.80	.031	271	520	.3540
Brass	8.51	.094			.3075
Cadmium	8.60	.057	321	610	.3107
Calcium	1.57	1.70	810	1490	.0567
Chromium	6.80	.120	1510	2750	.2457

PHYSICAL PROPERTIES OF METALS-Continued

			Meltin	z Point	Weight
	Specific	Specific	Deg. Cen-		in Lbs. per
Metal	Gravity	Heat	tigrade	renheit	Cubic Inch
Cobalt	8.50	.110	1490	2714	.3071
Copper	8.89	.094	1083	1982	.3212
Gold	19.32	.032	1063	1945	.6979
Iridium	22.42	.033	2300	4170	.8099
Iron	7.86	.110	1520	2768	.2634
T (C4)	# 010	1000	10==	2525	0.10
Iron (Cast)	7.218	.1298	1375	2507	.2605
Iron (Wrought)	7.70	.1138	1500-1600	2732-2912	.2779
Lead	11.37	.031	327	621	.4108
Lithium	0.57	.941	186	367	.0213
Magnesium	1.74	.250	651	1204	.0629
Manganese	8.00	.120	1225	2237	,2890
Mercury	13.59	.032	38.7	37.7	.4909
Monel Metal	8.87	.127	1360	2480	.320
Nickel	8 80	.130	1452	2646	.319
Platinum	21.50	.033	1755	3191	.7767
Potassium	0.87	1.70	62	144	.0314
Silver	10.53	.056	961	1761	.3805
Sodium	0.97	.290	97	207	.0350
Steel	7.858	.1175	1330-1378		.2839
Strontium	2.54	.074			.0918
Strouttum	2.04	.071	• • • •	••••	.0316
Tantalum	10.80		2850	5160	.3902
Tin	7.29	.056	232	450	.2634
Titanium	5.3	,130	1900	3450	.1915
Tungsten	19.10	.033	3000	5432	.6900
Uranium	18.70	• • • •	••••	• • • •	.6755
Vanadium	5.50		1730	3146	.1987
Zinc	7.19	.094	419	786	.2598
		-			

Protecting Aluminum from Corrosion Immerse for 10 minutes in bath of following at 50-60° C.

Formula No. 1

Sal Soda	125 g.
Sodium Chromate	8 g.
Ammonia	25 сс.
Water	1 l.

No. 2

Anodic treatment at 12 volts for 5 minutes and 15 volts for 5 minutes in following bath:

Oxalic .	Acid		25	g.
Sodium	Chromate		17	g.
Sodium	Dihydrogen	Sulphate	3	ğ.

Hardening Aluminum U. S. Patent 1,930,463

Pack in a mixture of:

fagnesium		95	11
Magnesium	Oxide	5	11

and heat at 420° C, in an atmosphere of carbon dioxide until the magnesium diffuses into the surface of the aluminum.

Non Serzing Aluminum U. S. Patent 1,978,112

Dip the aluminum in a bath of molten aluminum stearate.

Rustproofing Iron

U. S. Patent 1,949,921

Phosphoric Acid (8)	5%) 20 :	Ø,	OZ.
Ethyl Alcohol	20	fl.	oz.
Water	30	fl.	oz.
Isopropyl Ether	0.7-3.5	fl.	oz.

Radiator "Rust" Preventative U. S. Patent 1,940,041

Borax	36 lb.
Sodium Salicylate	30 lb.
Sodium Nitrate	7 lb.
Use 73 grains per quart	of water.

220	nr Chemica	D FORMUDARI	
Corrosion Inhi	hitae	from tools. If the solution	in unad man
		then one or two bound wi	is used warm,
Sodium Chromate	20 lb.	then one or two hours wi	ii sumce, but
Paraffin Oil	15 lb.	if used cold, it is best to s	llow the tools
Sulphonated Red Oil	50 lb.	to remain in the liquid	overnight. A
Liquid Soap	2 lb.	tablespoonful of the amm	onium citrat e
Soap Bark Extract	5 lb.	crystals may be used to a	pint of water,
Water to	make 100 lb.	although the proportions	are not im-
		portant. The solution w	rill serve re-
		peatedly until depleted.	
Non Corresive (Ethy	1) Alcohol	For tools of awkward a	hape such as
U. S. Patent 1,9	27,842	try squares and large ste	el sourres, n
About 0.03 per cent of		cardboard mailing containe	r may be used
ate or the equivalent of		in place of a vat, crock,	or other con-
have be the equivalent of	the correspond	tainer, if it is first impregi	nted with hot
borax, sodium lactate, or	allul to our	paraffin wax.	moca mon nou
ing potassium salts, is			
mercial alcohol to give pl			
venting corrosion of the n	ietal containers.	Rust and Oil Rem	over
		U. S. Patent 1,93.	5,911
Non-Corrosive Zinc Co	mdnit Alloy	Brush with:	
German Patent (314,996	Phosphoric Acid (75%)	69.5 lb.
621	02 05 1	Butyl "Cellosolve"	17 lb.
Zinc	83-95 kg.	Oleic Acid	0.5 lb.
Aluminum	13-3 kg.	Saponin	
Manganese	1 2 kg.	Water	1 lb.
Cadmium or Silicon	3-0 kg.	water	12 lb.
		Cleaning Motor Nan	ionlatos
Silver Tarnish Pr		Cleaning tarnish, grease	
British Patent -	130,795	the nameplate of motors a	and dirt ou
A jar containing the fol	lowing is placed	in order to read the figur	nu generators
in display cases containing		data is facilitated by the	es and other
Calcium Chloride,	8	data is facilitated by the of crinkled tin foil. The	use of a wad
Granular	88-94.9 g.	not scratched or marred by	minepiate in
	00-04.0 g.	as is the case when an ab	inis material
Copper Sulphate,	F 10	for removing the case when an an	rasive is used
Anhydrous	5-10 g.	for removing the accumula	tea airt.
Talc	0.1-2 g.		
Demoning Post fo	 Tuom	Decarbonizing Lining fo	r Cast Iron
Removing Rust fr		Molds	
Formula No.		Russian Patent 3	5,331
Soaking 12 hours in	Petroleum	Brown Iron Ore	68 lb.
No. 2		Refractory Clay	30 lb.
Make up:		Potassium Permanganate	2 lb.
	er		
Spindle Oil	65 g.	0.13 1 75 4 7	
Paraffin Scales or Ceres Yellow		Soldering Fluxes for Iro	
	15 g.	Ferrous Metal	8
Pumice Powder	20 g.	Stainless Steel	
No. 3		Borax	75-25 oz.
Dissolve:		Boric Acid	25-75 oz.
Water	1000 g.	Make into paste with alco	
Stannous Chloride	10 g.	Galvanized Iron	
Mercuric Chloride	2 g.	1	
N . 4	9.	Hydrochloric Acid	750 cc.
No. 4		Water	250 cc.
Use:		Zinc add until no more	will dissolve
Caustic Soda	10 g.	then add a solution of	•
Zinc Powder	10 σ.	A	

Removing Rust From Tools By using a solution of ammonium cit-rate, rust may be completely removed

Caustic Soda Zinc Powder

10 g. 10 g.

Ammonium Chloride Water

Stannous Chloride Water

then add following solution:

50 g. 170 cc.,

30 g. 170 cc.

Rosin Rosin Colore Rosin Ros	To form a paste solder of this type work in potato starch to desired consistency.	is stannous bromide 28, cadmium chlo- ride 20, cadmium iodide 10, animo- nium chloride 25, ammonium fluoride
Rosin Tallow, Ox 2 th. Zinc Chloride 1 lb. Tallow 40 lb. Rosin, Powdered 25 lb. Saturated Ammonium Chloride Solution 12½ lb. Tallow 2 lb. Olive Oil Saturated Ammonium Chloride Solution 2 lb. Saturated Ammonium Solder Formula No. 1 Tin 76 oz. Zins 20 oz. Altiminum Solder Formula No. 1 Tin 20 oz. Antimony 0.65 oz. Copper No. 2 French Patent 775,192 The solder contains cadmuna, lead and zine in the proportions of 4, 4, 3 and 2 lb./6 of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium 0.2 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zinconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum Soldering Fluxes British Patent 413,637 Copper 2 oz. Antimony one cation change characteristic standary in the usual way. Itard Solder for Cast Iron Copper 1 lb. Copper 2 lb. Disacteristic standary in the usual way. Itard Solder for Cast Iron Copper 2 lb. Copper 2 lb. Disacteristic standary in the usual way. Itard Solder for Cast Iron Soldering 1 lb. Copper 2 lb. Disacteristic standary in the usual way. Itard Solder for Cast Iron Copper 3 lb. Not	Aluminum Sheets	2, zinc chloride or zinc bromide 5%; 4
Tallow, Ox 2 1 h. Zine Chloride		paste with 6 parts of chlorodiphanet
Tallow, OX Zinc Chloride No. 2 Olive Oil No. 2 Olive Oil No. 2 Olive Oil Solution Tallow Aluminum Chloride Solution Aluminum Solder Formula No. 1 Tin Aluminum Solder Formula No. 1 Tin Zinc Aluminum Solder Formula No. 1 Tin Tin Zinc Aluminum Solder Formula No. 1 Tin Tin Zinc Aluminum 3 o.z. Aluminum 0.6 oz. Lead 0.2 oz. Copper No. 2 French Patent 775,192 The solder contains cadmum, lead and zinc in the proportions of 4, 4, 3 and 2-10% or zinc chloride. No. 3 French Patent 776,958 Zinc Soldering Iron Tip Alloy Bertish Patent 431,637 Copper 97 lb. Cobalt 2.6 lb. Beryllium 0.4 lb. Heat this for one hour at 900° C. Quench in water, reheat to 500° C. for one to two hours and allow to cool. Cast Iron Soldering Add to muritatic acid, zinc sufficient to ckill'' it, and drop in several small pacees copper before action censes. Use the twill be solution on cast iron that has been filed bright, and solder in the usual way. Hard Solder for Cast Iron Copper 60 lb. Tin 1 lb. Chain Link Solder U. S. Patent 2,003,865 Tin Copper 2 lb. Borax 3 lb. Hard Solders German Silver and Nickel Silver Copper 17 lb. Solver 65 lb. Copper 2 lb. Borax 3 lb. Tin 8 lb. Tin 8 lb. Tin 8 lb. Tin 8 lb. Copper 24 lb. Zinc 11 lb. Silver 30 lb. Copper 20 lb. Soldering Iron Tip Alloy Cobalt 2.6 lb. Copper 60 lb. Cast Iron Soldering Add to muritatic acid, zinc sufficient to ckill'' it, and drop in several small pacees copper before action censes. Use the ckill' it, and drop in several small pacees copper before action censes. Use the ckill' it, and drop in several small pacees copper before action censes. Use the ckill' it, and drop in several small pacees copper before action censes. Use the ckill' it, and drop in several small pacees copper before action censes. Use the ckill' it, and drop in several small pacees copper before action censes. Use the ckill' it, and drop in seve		
Olive Oil Olive Oil Tallow Aluminum Solder Formula No. 1 Tin Aluminum Solder Formula No. 1 Tin Tin Aluminum Solder Formula No. 1 Tin Tin Aluminum Solder Formula No. 1 Tin Aluminum Solder Formula No. 1 Tin Tin Aluminum Solder Formula No. 1 Tin Aluminum Solder Formula No. 1 Tin Tin Tin Tin Tin Tin Tin Tin Tin Ti		- Tarana de la caración de la caraci
Olive Oil Tallow Rosin, Powdered Solution Tile Rosin, Powdered Tile Rosin Rosin, Powdered Tile Rosin, Powdered Tile Rosin, Powdered Tile Rosin Water, reheat to 500° C, for one to two hours and allow to cool. Cast Iron Soldering Add to muriatic acid, zinc sufficient to "kill" it, and drop in several small process copper before action ceases. Use this solution on east iron that has been filed bught, and solder in the usual way. Ital Solder for Cast Iron Copper Roper Rosill' it, and drop in several small process copper before action ceases. Use this solution on east iron that has been filed bught, and solder in the usual way. Ital Solder for Cast Iron Copper Roper Rosill' it, and drop in several small process copper before action ceases. Use this solution on east iron that has been filed bught, and solder in the usual way. Ital Solder for Cast Iron Copper Roper Rosill' it, and drop in several small process copper before action ceases. Use this solution on east iron that has been filed bught, and solder in the usual way. Ital Solder for Cast Iron Copper Rosill' it, and drop in several small Casin Is and solder in the usual way. Ital this for one hour at loof of the 'kill' it, and drop in several small Cast Iron Soldering Capper Rosill' it, and to muriatic acid, zinc s	Zinc Chloride 1 lb.	C.11 T Mi All.
Olive Oil 50 lb. Tallow 40 lb. Rosin, Powdered 25 lb. Saturated Ammonium Chloride Solution 12½ lb. Tin Resin, Powdered 1 lb. Tallow 2 lb. Tallow 2 lb. Saturated Ammonium Chloride Solution 2 lb. Tallow 2 lb. Saturated Ammonium Chloride Solution 2 lb. Saturated Ammonium Solder Formula No. 1 Saturated Ammonium Solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before action ceases. Use this solution on cast iron that has been filed linght, and drop in several small pieces copper before	No. 2	
Tallow 40 lb. Rosin, Powdered 25 lb. Saturated Aumonium Chloride Solution Tide Solution Tine Rosin, Powdered 1 lb. Tallow 2 lb. Olive Oil 2 lb. Saturated Ammonium Chloride Solution Aluminum Solder Formula No. 1 Tin 76 oz. Zine 20 oz. Aluminum 3 oz. Antimony 0.6 oz. Lead 0.2 oz. Copper 0.2 oz. Antimony 0.6 oz. Lead 0.2 oz. Copper 0.2 oz. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 lb/Silicon 0.4 lb. Iron 0.4 lb. Silicon 0		British Patent 431,637
Resin, Powdered 1 lb. Tallow 2 lb. Saturated Ammonium Chloride Solution 12½ lb. Tallow 2 lb. Saturated Ammonium Chloride Solution 2 lb. Aluminum Solder Formula No. 1 Tin 76 oz. Zine 20 oz. Aluminum 3 oz. Antimory 0.5 oz. Lead 0.2 oz. Copper 0.2 oz. Copper No. 2 French Patent 775,192 The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 logs of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Silicon 0.		Copper 97 lb.
ride Solution 12½ lb. Tin Rosin, Powdered 1 lb. Tallow 2 lb. Olive Oil 2 lb. Saturated Ammonium Chlorde Solution 2 lb. Aluminum Solder Formula No. 1 Tin 76 oz. Zino 20 oz. Aluminum 3 oz. Aluminum 3 oz. Copper 0.2 oz. Copper 0.2 oz. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 lb/s of zinc chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zirconium No. 4 British Patent 426,526 Zine 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iolide, ammonium chloride, zinc chloride, zinc bloride, zinc bl	Rosin, Powdered 25 lb.	
Tin Rosin, Powdered 1 lb. Tallow 2 lb. Colive Oil Saturated Ammonium Chlorides Pormula No. 1 Tin 76 oz. Zime 20 oz. Aluminum 3 oz. Antimony 0.6 oz. Lead 02 oz. Copper 0.2 oz. Copper 0.2 oz. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 lb/5 of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zirconium No. 4 British Patent 426,526 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zirconium 0.2 lb. Mercury 3 lb. Aluminum 11/4-1 lb. Lead 1/4-1 lb. Aluminum 11/4-1 lb. Aluminum 11/4-1 lb. Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc bromide fluorides, cliorodiphenyl, p-di- Bornal Silver 1 lb. Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodiphenyl, p-di- Bornal Silver 30 lb. Cust Iron Soldering to cust Iron Soldering to chill to chill' it, and drop in several small pacces copper hefore action ceases. Use thins solution on cast iron that has been filed hight, and solder in the usual way. Hard Solder for Cast Iron Capper 60 lb. Tim 1 lb. Chain Link Solder U. S. Patent 2,003,865 Tim 1 lb. Copper 2 lb. Blorax 3 lb. Hard Solders German Silver and Nickel Silver 65 lb. Copper 2 lb. Silver 65 lb. Copper 24 lb. Zinc 11 lb. Capper 24 lb. Zinc 11 lb. Copper 50-60 lb. Nickel 3 lb. Zinc 37-27 lb. Soft Soldering Monel and Nickel Monel metal and nickel are soft	Saturated Ammonium Chlo-	Beryllium 0.4 lb.
Rosin, Powdered Tallow Olive Oil Saturated Ammonium Chloride Solution Aluminum Solder Formula No. 1 Tin 20 0z. Aluminum 3 0z. Antimony 0.5 0z. Lead 0.2 0z. Copper 0.2 0z. Copper 0.2 0z. Copper No. 2 French Patent 775,192 The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2: 10% of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Silicon 0.5 lb. Tin 14 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bromide fluorides, chlorodiphenyl, p-di- Bornax 3 lb. Soft Soldering Monel and Nickel Soft Soldering Monel and Nickel Monel metal and nickel are soft	ride Solution 121/2 lb.	Heat this for one hour at 900° C.
Tallow Olive Oil Saturated Ammonium Chlorder Solution Aluminum Solder Formula No. 1 Tin 76 oz. Zine 20 oz. Aluminum 3 oz. Antimony 0.6 oz. Lead 0.2 oz. Copper 0.2 oz. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 logs of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zirconium No. 4 British Patent 426,526 Zine 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Lead 1½-1 lb. Lead 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iolide, ammonium chloride, zinc chloride, zinc bloroide, zinc chloride, zinc bloroide, ammonium chloride, zinc chloride, zinc bloroide, zinc bloroid	Tin	Quench in water, reheat to 500° C. for
Tallow Olive Oil Saturated Ammonium Chloride Solution 2 lb. Saturated Ammonium Chloride, aim one of the following. Eadminum Solder Formula No. 1 Tin 76 oz. Zino 20 oz. Aluminum 3 oz. Antimony 0.6 oz. Lead 0.2 oz. Copper 0.2 oz. Copper No. 2 French Patent 775,192 The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 lb/6 of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Iron 0.4 lb. Iron 0.4 lb. Iron 0.4 lb. Silicon 0.4 lb. Silicon 0.4 lb. Iron 0.2 lb. Tin 1 lb. Copper 2 lb. Hard Solders British Patent 426,526 Zine 22 lb. Tin 14 lb. Mercury 3 lb. Mercury 3 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Lead ½-1 lb. Lead 1/½-1 lb. Lead 1/½-1 lb. Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bromide fluorides, chlorodiphenyl, p-di- Solver 3 lb. Cnst Iron Soldering soil, zine sufficient to ckill' it, and drop in several small porces copper before action ceases. Use this solution on east iron that has been filed lunght, and solder in the usual way. Hard Solder for Cast Iron Copper 40 lb. Tin 1 lb. Chain Link Solder U. S. Patent 2,003,865 Tin 1 lb. Copper 2 lb. Silver 3 lb. Tin 8 lb. Tin 9 lb. Copper 3 lb. Copper 3 lb. Silver 3 lb. Si	Rosin, Powdered 1 lb.	one to two hours and allow to cool.
Olive Oil Saturated Ammonium Chlorde Solution Aluminum Solder Formula No. 1 Tin 76 oz. Zine 20 oz. Aluminum 3 oz. Antimony 0.6 oz. Lead 02 oz. Copper 0.2 oz. Copper 0.2 oz. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 liv/s of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Zirconium 0.2 lb. No. 4 British Patent 426,526 Zine 22 lb. Tin 14 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc bromide fluorides, chlorodichenyl, p-di- Formula No. 1 Cantal Iron Solder in the usual way. Hard Solder for Cast Iron Copper 60 lb. Zine 40 lb. Tin 1 lb. Copper 2 lb. Borax 3 lb. Hard Solders Tin 8 lb. Tin 14 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodichenyl, p-di- Soft Soldering Monel and Nickel Monel metal and nickel are soft		
Saturated Ammonium Chlorides and solution 2 lb. Aluminum Solder Formula No. 1 Tin 20 02. Zina 20 02. Aluminum 3 0.5 02. Antimony 0.6 02. Lead 0.2 02. Copper 0.2 02. Copper 0.2 02. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2 lb/6 of zine chloride. No. 3 French Patent 776,958 Zine No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Silicon 0.4 lb. Silicon 0.4 lb. Silicon 0.4 lb. Sireonium 0.2 lb. No. 4 British Patent 426,526 Zine 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bromide, floorides, chlordide, long collection, and not collected and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bromide flourides, chlorodiphenyl, p-di- Monel metal and nickel are soft	Olive Oil 2 lb.	Cast Iron Soldering
Tin	Saturated Ammonium Chlo-	Add to muriatic acid, zine sufficient to
Aluminum Solder Formula No. 1 Tin	ride Solution 2 lb.	"kill" it, and drop in several small
Aluminum Solder Formula No. 1 Tin 76 oz. Zino 20 oz. Aluminum 3 oz. Antimony 0.6 oz. Lead 0.2 oz. Copper 0.2 oz. Copper No. 2 French Patent 775,192 The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2: 10% of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Silicon 0.4 lb. Silicon 0.4 lb. Zirconium 0.2 lb. Tin 14 lb. Aluminum 124-1 lb. Aluminum 124-1 lb. Lead 22 lb. Tin 14 lb. Lead 32 lb. Aluminum 124-1 lb. Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bromide, fluorides, chlorodiphenyl, p-di- Borniad Solder for Cast Iron Copper 60 lb. Tin 1 lb. Chain Link Solder U. S. Patent 2,003,865 Tin 1 lb. Copper 2 lb. Borax 3 lb. Hard Solders U. S. Patent 2,003,865 Tin 1 lb. Copper 2 lb. Borax 3 lb. Tin 8 lb. Tin 22 lb. Copper 50-60 lb. Nickel 3 lb. Zinc 37-27 lb. Soft Soldering Monel and Nickel Monel metal and nickel are soft		pieces copper before action ceases. Use
Formula No. 1 Tin	Alaminan Galdon	
Tin		
Hard Solder for Cast from Aluminum		***************************************
Antimory		Hard Solder for Cast Iron
Antimony 0.6 oz. Lead 0.2 oz. Copper 0.2 oz. Copper 0.2 oz. The solder contains cadmum, lead and zine in the proportions of 4, 4, 3 and 2-10% of zine chloride. No. 3 French Patent 776,958 Zine 89-95 lb. Aluminum 10-4 lb. Silicon 0.4 lb. Iron 0.4 lb. Iron 0.4 lb. Zirconium No. 4 British Patent 426,526 Zine 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Lead 1½-1 lb. Lead 1½-1 lb. Lead 1½-1 lb. Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iolide, ammonium chloride, zinc chloride, zinc bromide fluorides, chlorodichenyl, p-di-formide fluorides, chlorodichenyl fluorides, chlorodichenyl fluorides, chlorodichenyl fluoride, chlorodichenyl fluori		1
Tim		
Copper		
No. 2		
Trench Patent 775,192 Chain Link Solder U. S. Patent 2,003,865		
U. S. Patent 2,003,865		(1) 1 211 (2)
Time in the proportions of 4, 4, 3 and 2	_	
No. 3	The solder contains cadmum, lead and	U. S. Patent 2,003,865
No. 3 Borax 3 lb.	zine in the proportions of 4, 4, 3 and 2-	Tin 1 1b.
French Patent 776,958 Zine	10% of zinc chloride.	
Zine	No. 3	Borax 3 lb.
Zine	French Patent 776,958	Mari Caller
Aluminum		1
Silicon		
Tron		
No. 4 British Patent 426,526 Zinc 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc bloromide, fluorides, chlorodic, pdi- Silver 30 lb. Clopper 40-50 lb. Zinc 20-30 lb. Austentic Stainless Steels Silver 30 lb. Zinc 20-30 lb. Nickel 3 lb. Zinc 37-27 lb. Soft Soldering Monel and Nickel Monel metal and nickel are soft	Iron 0.4 lb.	
British Patent 426,526 Zinc 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc bloromide fluorides, chlorodiphenyl, p-di- Soft Soldering Monel and Nickel Monel metal and nickel are soft	Zirconium 0.2 lb.	
Zine 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bromide fluorides, chlorodiphenyl, p-di- Soft Soldering Monel and Nickel Monel metal and nickel are soft	No. 4	Thin Copper
Zine 22 lb. Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine bloromide fluorides, chlorodip. Pd. Pd. Soft Soldering Monel and Nickel Monel metal and nickel are soft	British Patent 426,526	
Tin 14 lb. Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium ioidide, ammonium chloride, zinc chloride, zinc bloroide, elhoroide, elhoroide		
Aluminum 142-1 lb. Lead 42-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local control of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, zinc chloride, zinc chlori		
Lead 1/2-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, pulsonium chloride, zinc chloride, zinc blorodide, logical standard	110.	Zine 11 lb.
Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, horoides, chlorodide, pdi- Soft Soldering Monel and Nickel Monel metal and nickel are soft	Mercury 3 lb.	
Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodichenyl, p-di- bromide fluorides, chlorodichenyl, p-di- Monel metal and nickel are soft	Mercury 3 lb. Aluminum 1½-1 lb.	Henvy Brass Silver 30 lb.
British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, blorodide, chlorodide, pdi- British Patent 413,141 Silver 10 lb. Copper 50-60 lb. Nickel 3 lb. Zinc 37-27 lb. Soft Soldering Monel and Nickel Monel metal and nickel are soft	Mercury 3 lb. Aluminum 1½-1 lb.	Heavy Brass Silver 30 lb. Copper 40-50 lb.
Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, local promide fluorides, chlorodide, pdi-	Mercury 3 lb. Aluminum 1½-1 lb. Lend ½-1 lb.	Heavy Brass Silver 30 lb. Copper 40-50 lb.
Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, chlorodide, pdi-blorodide, chlorodide,	Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes	Henvy Brass Silver 30 lb. Copper 40-50 lb. Zinc 20-30 lb.
cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodide, chlorodide, pdi-bromide fluorides, chlorodide, pdi-bromide, pdi-bromide	Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes	Heavy Brass 30 lb.
preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc bromide fluorides, chlorodiphenyl, p-di-	Mercury 3 lb.	Heavy Brass 30 lb. Copper 40-50 lb. Zinc 20-30 lb. Austentic Stainless Steels Bilver 50-60 lb. Copper 50-60 lb.
one of the following: cadmium iodide, ammonium chloride, zinc chloride, zinc blorodio bromide fluorides chlorodio benyl, p-di- Monel metal and nickel are soft	Mercury 3 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing	Heavy Brass 30 lb.
bromide fluorides, chlorodiphenyl, p-di- Monel metal and nickel are soft	Mercury 3 lb. Aluminum 1½-1 lb. Lead ½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than	Heavy Brass 30 lb.
chlorbenzene. A preferred composition soldered readily. Many of the soft	Mercury 3 lb. Aluminum 1½-1 lb. Lead 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide,	Heavy Brass 30 lb.
emiordenzene. A preferred composition i somered readily. Many of the soft	Mercury Aluminum 1½-1 lb. Lead 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride zinc chloride, zinc	Heavy Brass Silver 30 lb.
	Mercury Aluminum 1½-1 lb. Lead 1½-1 lb. 1½-1 lb. Aluminum Soldering Fluxes British Patent 413,141 Claim is made for fluxes containing cadmium chloride and stannous bromide, preferably in admixture with more than one of the following: cadmium iodide, ammonium chloride, zine chloride, zine promide fluoride, zine chloride, zine promide fluoride, zine chloride, zine	Heavy Brass 30 lb.

solders regularly used in the copper shop will make suitable joints in both of these metals. In making a lock seamed joint, for example, it is definitely recommended that the edges of the sheet be tinned, that is, coated with a thin film of soft solder before forming the sheet and before lock seaming.

Once the sheet has been properly tinned, it is then very easy to flow in the soft solder and make a tight joint which is reasonably strong.

Similarly, in sweating a tube into a header, if both the header and end of tube are tinned first, then assembled, heated, and solder flowed in, a sound joint will be obtained. It is necessary in all soldering work to have surfaces clean and bright if joints are to hold.

It must be remembered that the strength and ductility of soft solder is not of a very high order and for that reason soft solder is not recommended where considerable vibration is apt to be involved.

Soldering Flux for Stainless Steel U. S. Patent 1.968,841

Boric acid, three parts; borax, two to three parts; and ammonium chloride, one and one-half to three parts, together with a liquid from the group consisting of water and hydrogen peroxide, in quantity to make a thick pasto.

Soldering Flux for Stainless	St	eel
Zinc Chloride	37	oz.
Acetic Acid	23	oz.
Hydrochloric Acid	40	oz.
Name of the last o		

Tin Plate Sold	ler
Ammonium Chloride	4 oz.
Zinc Chloride	48 oz.
Hydrochloric Acid	1 oz.
Water	47 oz.
Dilute to required streng	th with water.

Crankshaft Heat Treatment

Shafts are heat treated in gas fired furnaces as follows:

Heat to 1650° F., hold for 20 minutes. Air quench to a minimum of 1200° F. Reheat to 1480° F., and hold 1 hour. Cool in the furnace to 1000° F. in another hour.

The alloy for casting is melted in four 15-ton electric furnaces according to the latest approved practice. The charge is

made up of approximately 50 per cent return shop scrap (gates and risers) and 50 per cent steel scrap.

Drawhead Casting Heat Treatment

To obtain the best combination of mechanical properties, the castings are given a simple heat treatment, as follows:

Heat to about 1650° F., hold at heat 1 to 1½ hours per inch thickness of heaviest section, and cool in still air to a black heat. Reheat to 1200-1250° F., hold at least 1 hour per inch and cool in air or furnace. This treatment is not difficult and can be performed very readily with ordinary equipment. Usually the cost of such a treatment is no greater than that for simple annealing.

Oil Well Tool Heat Treatment

The heat treatment of these steels (used either for slip socket or tool joint) is quite similar. After forging it is advisable to anneal the steel to relieve any forging strains and at the same time put it in a readily machinable condition. One of the simplest treatments for doing this is to heat the steel to above 1600/1650° F. and cool in the furnace until black or, if removed from the furnace, pack in lime or ashes so that it cools slowly. The forging should then be machined and the final heat treatment performed as follows:

Heat to about 1550° F.; hold at this temperature until heated through thoroughly; quench in oil. The tempering operation will depend upon the hardness specifications. This steel is quite tough in the hardness range of 280/320 Brinell which could be secured by using a drawing temperature around 900° F, holding at this temperature until heated through thoroughly in the heavy sections. While final machining can be performed in this hardness range, it must be done very slowly, and it is desirable to use a lower hardness range, such as about 240/280 Brinell which is obtained with about 550° F. draw. The physical properties secured at these hardnesses is about as follows:

Tensile Strength	145,000 p.s.i.
Yield Point	120,000 p.s.i.
Elongation in 2"	18%
Reduction of Area	57%

For the slips a case hardened steel such as S.A.E. 2315 is used, arrangements to be made so that only the teeth are case hardened. This can be accomplished

by copper plating the piece before cutting the teeth so that the copper remains on all the parts except the teeth. Use a case hardening temperature of 1650/ 1700° F., cooling in the box and re-heating the parts to a temperature of 1475° F., quenching in oil, and tempering by heating to 275-300° F. This treatment will toughen the core of the part so that it will be sufficiently hard not to stick or gall against the socket.

Heat Treatment of High Strength Shafting

Heat treatment for S.A.E. 3340: Oil quench from 1500° F. and temper at 800° to 900° F.

Heat treatment for Ni-Cr-Mo: Oil quench from 1575° F., temper at 900° to inone F

Brake Drum Heat Treatment

The heat treatment given brake drums is heating to 1600° F., holding there for 30 min., then cooling rapidly in the furnace to 1450° F., followed by cooling in 2 hours to 1350° F. and then in 1 hour to 1000° F.

Valve Gear Metal Heat Treatment

A nickel-molybdenum case hardening steel corresponding to S.A.E. composition 4615 is used. This material can be machined to the finished size in a soft state and then should be carburized by the pack method, at a temperature between 1650 and 1700° F., until a case about 1/32 in. in depth is secured. For the best results we would recommend quenching from the carburizing box into oil. This should be followed by a reheating to a temperature of 1375 to 1400° F. and quenching in oil, then temper at about 275° F. This treatment will result in a very hard case which should show excellent wearing properties.

Carburizing Nickel Steel

(1) A simple and economical treatment where refinement of the case is not important, is to carburize at 1600° F. and quench in oil directly from the box, followed by tempering at 250 to 350° F. (2) Or, if cooled in the box after carburizing, then heat to 1475-1500° F. and oil quench, then temper as above, to get a refined and tough core which will back up the hardness of the case. (This is not recommended if the carbon content of the core is over about .18%, as brittleness may result.)
(3) Cooling in the box, oil quenching

from 1325 to 1375° F. and tempering, is recommended where a hard and refined case is the main requirement. (4) If refinement of both case and core is demanded, and economy and speed is not so important, a double treatment should be given, as follows: Carburize at 1600° F., cool in box. Quench in oil from 1500-1550° F., and again from 1325-1375°. Temper at 250-350° F. as required. This will give a very hard case and a ductile core, and is much used on gears of fine pitch.

Grinding Wheels U. S. Patent 1,937,043

Carborundum 900 g., is mixed with furfuraldehyde 10 cc. till moist then with a phenolic resin 100 g., and the mixture is pressed into shape at less than 80° C. The articles are then heated at a suitable temperature until complete hardening occurs.

Aluminum Welding Flux

Aid	minam werang	I IUA
Potassiur	n Chloride	79 oz.
Salt		16 oz.
Potassiur	n Bisulphate	5 oz.
	re is best used	
aluminum c	ontaining 4% si	licon.

Bronze-Welding

Bronze welding, as a general term for actual bronze-welding and for bronzesurfacing, is used today for joining metals of high melting points, as cast non, steel, nickel, copper and their alloys, by the use of a bronze bonding material. For use with the oxy-acetylene flame, rod of 59% copper, 40% zing od 35, is generally used, while recently other elements as silicon, manganese, from kg. been added. Lead is objectionable increases porosity of the weld metal.

Welding Rods for Copper, Steel and Bronze

U. S. Patent 2,009,977

	,	•		
Silicon			3.5	lb.
Tin			0.5	lb.
Phosphorus			0.05	lb.
Copper			96	lb.

Welding Zinc and Zinc Alloy Castings The welding of zinc requires some care because of its low melting point and the tenacious character of the oxide. A gas flame should be used with welding rod of the same metal and a flux of ammonium chloride and water. The welding operation always weakens the surrounding metal and should, if possible, be followed by a cold working operation to refine the grain.

Zinc alloy castings containing aluminum are extremely difficult to weld and the success of the operation depends largely on the technique of the welder.

Welding Electrode Coating Canadian Patent 341,572 Formula No. 1

Shredded wood 100, sodium silicate 80, calcium carbonate 5, kaolin 5, silicomanganese 5 and peanut oil 5 parts. The coating in a plastic state is applied to the core and then baked or dried.

No. 2

U. S. Patent 1,968,984

Barium Chloride 20-50 lb.
Lithium Fluoride 4-6 lb.
To the above add 75-45% of following mixture.

Salt 40-50 lb. Potassium Chloride 60-50 lb.

No. 3

U. S. Patent 2,000,861

Slip clay 40-60 parts, iron oxide 20-30 parts, calcium carbonato 20-30 parts, feldspar 15-30 parts, rutile 5-20 parts, manganese ore 5-15 parts, carbonaceous material 5-15 parts, ferromanganese 5-20 parts, ferrocheme 2-8 parts and dextrin 1-15 parts by weight.

Welding Rod for Bearing Metals U. S. Patent 1,926,412

Zinc	90 1	b.
Copper	5 1	b.
Antimony	5 1	b.

Welding Rod Conting Formula No. 1

Canadian Patent 347,320

Suspend above in sufficient ion of	of a	solu-
Calcium Fluoride		lb.
Titanium Dioxide	22	lb.
Barium Carbonate	9	lb.
Calcium Carbonate	8	ID.

Potassium Silicate 2 lb. Water No. 2

U. S. Patent 1,992,792

Titanium Dioxide	1 lb.
Tale	1 lb.
Feldspar	1 lb.
Sodium Silicate	3 lb.
Water	to suit

Aircraft Engine Alloys

Use case hardened 5% nickel steel (S.A.E. No. 2512) for aircraft engine gears. The crankshafts should be forged of a nickel-chromium steel such as S.A.E. 3240, or nickel-chromium-molybdenum steel of the following approximate composition:

errion:	
Carbon	0.40-0.50 lb.
Manganese	0.45-0.75 lb.
Nickel	1.50-2.00 lb.
Chromium	0.60-0.90 lb.
Molybdenum	0.15-0.25 lb.
Iron	to make 100 lb

Heavy Duty Axle Alloy

Carbon	0.35-0.45 lb.
Nickel	1.50-2.00 lb.
Chromium	0.60-0.80 lb.
Manganeso	0.60-0.80 lb.
Molybdenum	0.30-0.40 lb.
Iron	to make 100 lb.

Nickel Steel Pin and Bearing Alloy

5% nickel steel such as S.A.E. 2512, with the carbon at the upper end of the range, say 0.15% is used.

Carburize this steel at 1600-1650° F. The most suitable depth of case will depend upon the dimensions of the pin, and normally should not be more than 15% of its diameter. The cooling after carburizing should preferably be done in the box, but it is recommended that it be as rapid as convenient, such as allowing the box to cool in free air or possibly in an air blast.

For the hardening operation a single quench would be advisable at a temperature just high enough to refine the core. On these small pieces a temperature around 1440-1450° F. would be sufficient. The tempering operation on this steel should be at 275° F. The complete treatment will give maximum core strength, combined with very good toughness.

Hard Tool Steel Alloys Japanese Patent 101,748

Mold following under high pressure at 1600-1800° C.

Formula No. 1

Vanadium Powder	5 lb.
Tungsten Carbide	95 lb.
No. 2	

Titanium Powder 5 lb. Tungsten Carbide 95 lb.

No. 3	
Vanadium	3 lb.
Titanium	2 lb.
Tungsten Carbide	95 lb.

Steering Knuckle and Spring Bolt Alloy
A case hardened steel of the following
composition is used.

Carbon	0.12-0.20 lb.
Manganese	0.30-0.60 lb.
Nickel	3.25-3.75 lb.
Molybdenum	0.20-0.30 lb.
Iron	to make 100 lb.

Punch and Die Alloys

Use steels containing

 Carbon
 0.6–0.65
 lb.
 0.6–0.65
 lb.

 Manganese
 0.3–0.6
 lb.
 0.3–0.6
 lb.

 Nickel
 1.5–2
 lb.
 0.5–0.8
 lb.

 Chromium
 0.9–1.25
 lb.
 0.6–0.8
 lb.

 Molybdenum
 —
 0.2–0.4
 lb.

 Iron
 to make 100
 lb.

It should be thoroughly annealed after forging as follows: Heat to 1550/1575° F., air-cool, reheat to 1200/1250° F., hold for 6 to 8 hours and cool very slowly. To harden, heat to 1425° F., quench in oil, and temper for 1 hour at 425/450° F.

Shovel Dipper Teeth Alloy

puovei Dipper	reeth Anoy
Carbon	0.4-0.50 lb.
Nickel	3.0-3.50 lb.
Chromium	1.0-1.25 lb.
Molybdenum	0.3-0.40 lb.
Iron	to make 100 lb.
Heat treatment:	

Heat to 1750° F., hold 11/2 hours per

inch thickness; air cool. Reheat to 1250° F., hold at least one-hour per inen thickness; cool in air or furnace. Some foundries furnish the teeth in this condition, while others claim better wear by giving the tips a second treatment for hardening. This is done by heating the point or tip to a distance of 2 in. or 3 in. (dependent upon the size and shape of the tooth), to a red heat (1500-1800° F.) and cooling rapidly with an air blast. If it is found that the points are too brittle the whole tooth may be drawn at 700-800° F. Sometimes this tip hardening treatment is given to the castings after a plain annealing of the whole tooth, thus eliminating original air quenching and drawing treatment described at the beginning of this paragraph.

Another steel used quite successfully for shovel teeth in this service is the following:

Carbon	0.40-0.5 lb.	
Nickel	1.75-2.0 lb.	

Chromium	0.70-0.9	lb.
Iron	to make 100	lb.

These castings are given either an annealing or air quenching treatment as described above for the nickel-chromium-molybdenum steel. The tips are then reheated to a red heat and quenched in oil. The whole casting is then drawn at 700-900° F., depending upon the hardness required.

A steel which is giving excellent service in custings subjected to wear, has the following composition:

nowing composition.	i
Carbon	0.35-0.45 lb.
Manganese	1.25-1.50 lb.
Nickel	2.25-2.50 lb.
lron	to make 100 lb.

Acid Resisting Alloy

Patented

Molybdenum	0.5-10	lb.
Tin	4-5	lb.
Lead	95.5-85	lb.

Antifriction Alloy British Patent 413,209

Copper	67.5 lb.
Leid	25 lb.
Tm	5 lb.
Nickel	1 lb.
Antimony	0.5 lb.
Cadmium	0.5 lb.
Zinc	0.5 lb.

Hard Aluminum Alloy British Patent 406,161

31.09402	
89 -94 lb.	
1.5 lb.	
3.7- 5.5 lb.	
0.2- 1 lb.	
0.2 - 1 lb.	
0.4- 2 lb.	
	89 -94 lb. 1.5 lb. 3.7- 5.5 lb. 0.2- 1 lb. 0.2- 1 lb.

Aluminum Alloy for Chill Casting U. S. Patent 1,997,494

C. D. Tatent	1,001,101	
Aluminum	75 -95	lb.
Iron	2 -10	lb.
Antimony	0.5-15	lb.
Magnesium	0.2- 0.4	lb.

Oxidizing Nickel Silver

CARDINING THEACT	Direct	
Hydrochloric Acid	1	gal.
White Arsenic	10	oz.
Copper Sulphate	10	oz.
Ferric Chloride	2	OZ.
Copper Acetate	2	0 z.
Ammonium Chloride	1	OZ.
Hyposulphate of Soda	11/4	0 2.

Heat the hydrochloric acid, and when hot put in the white arsenic. When the white arsenic is completely dissolved, mix in the balance of the formula.

It must be definitely understood that this solution can only be used while cold. The article can be placed in a plater's basket or wired and dipped possibly half a dozen times in the solution, rinsed in cold water and then dipped in a solution of sodium cyanide and then rinsed again in cold water. After this rinse, the article should again be dipped in the

oxidizing solution and the process is then complete.

The result should be a jet black oxide which can be scratch brushed if a solid black is desired, and can be readily spotted off for highlights.

Copper Alloy Resistant to Sea Water U. S. Patent 1,956,251

Bilicon	1 - 3.25	lb.
Tin	0.5 - 1.5	lb.
Iron	0.75- 1.27	lb.
Lead		lb.
Copper	97.75-93.98	

Copper Alloy Spot Welding Electrode U. S. Patent 1,957,214

Cold Working Copper Alloy U. S. Patent 1,936,397

 Silicon
 0.75 lb.

 Manganese
 0.25 lb.

Non-Staining Copper Alloy U. S. Patent 2,007,430

Nickel	1 to 5	lb.
Cobalt	0.25 to 2	lb.
Silicon	0.25 to 2	lb.
Aluminum	1 to 5	lb.
Molybdenum	0.25 to 3	lb.
Iron	0.10 to 1	lb.
Calcium	0.05 to 0.5	lb.
Copper of an	amount to compl	ete
100 lb. mass.		

High Melting Copper Alloy German Patent 597,938

Beryllium	0.3-10 lb.
Aluminum	0.5-12 lb.
Copper	99.2–88 lb.

L	o₩	Cost	Dental	Alloy
---	----	------	--------	-------

Silver	85 oz.
Gold	10 oz.
Palladium	5 oz.

Cheap Dental Inlay Alloy

Oncar	Dentar	Imiay	Alloy	
Copper			19.29	lb.
Silver			79.29	lb.
Zinc			0.71	lb.
Tin			0.71	lb.

Cast Denture Alloy Canadian Patent 342.946

Chromium	17.5 lb.
Cobalt	57 lb.
Tungsten	3 lb.
Nickel	21 lb.
Iron	1 lb.
Carbon	0.5 lb.

Dental Alloy

	1 I CHOIL	I atent	20,121	
Gold			20-15	oz,
Copper	•		3-12	oz.
Silver			65-63	oz.
Zinc			7-8	oz.

Dental Filling Alloy German Patent 603,456

Bismuth	62.5 g.
Tin	37.2 g.
Gallium	1.3 g.

Dental Alloy Casting Mold British Patent 412,303

Plaster of Paris	40	lb.
Cristobalite	45	lb.
Tridymite	10	lb.
Quartz	5	lb.

Dental and Jewelry Alloy U. S. Patent 1,965,012

Gold	5-15	0 Z .
Palladium	22-30	0 Z .
Bilver	37-50	oz.
Copper	10-20	0 z .
Indium	0.5-5	

Imitation Gold Alloy French Patent 776,806

Copper	80-82 g.
Zinc	11-15 g.
Tin	3–5 g.
Nickel	2 g.

During fusion add the following per 100 g. of alloy. Cream of Tartar Magnesium Oxide 6 g. 3.5 g. Ammonium Chloride 'Lime 1.5 g.

Lead Calcium Alloys British Patent 412,316

Lead and pea size pieces of calcium carbide are mixed at 650-700° C. in presence of fused slag consisting of salt, calcium chloride and calcium fluoride. Alloys containing 3-3.5% calcium are obtained in 8 to 10 hours.

Lead Storage Battery Alloy British Patent 411,524

Tellurium	0.05	lb.
Antimony	6	lb.
Lead	93.95	lb.

Non-Corrosive Magnesium Alloy German Patent 613,511

Zinc	1-10	lb.
Iron	0.02 - 1	lb.
Silver	0.05 - 3	lb.
Magnesium	98.93-86	lb.
_		

Radium Beam Therapy Alloy Nickel 5 lb. 5 lb. Copper 90 lb. Tungsten Sinter the powdered metals at 1250-

> Arc-Light Reflector Alloy German Patent 615,119

1350° C.

Cobalt or Nickel 20-60 lb. Tungsten or Molybdenum 15-50 lb. 30-40 lb. Chromium 1-5 lb. Carbon or Silicon

Electric Light Reflector Alloy British Patent 412,074

Mirrors of Silver-Copper Alloy Canadian Patent 348,131

Prepare solution No. 1 by adding to 16 oz. of silver nitrate, 11 oz. of ammonia (26°) and, after the solution is complete, 16 oz. of distilled water; cool,

filter and add to the filtered solution an additional 144 oz. of distilled water.

Prepare solution No. 2 by dissolving 1 lb. crystallin copper sulphate in 64 oz. of distilled water, filter and place in a dark bottle.

For solution No. 3, to 64 oz. of distilled water add 2 lb. of crystallin Rochelle salt, heat to boiling and add 1 oz. of silver nitrate dissolved in 4 oz. of distilled water. To this mixture at the boiling point add 4 oz. of solution No. 2 and boil for at least 10 minutes; then cool, filter and place the filtered solution in a dark bottle.

For solution No. 4, dissolve 1 lb. of powdered tartaric acid in 48 oz. of distilled water, let stand 1 week and filter.

Prepare the final solution from distilled water, 64 oz.; solution No. 1, 2 oz.; solution No. 3, 2 oz.; and solution No. 4, 3 dr. Polish and brush with water the glass that is to be coated; then apply a weak solution of tin chloride with a felt block or bristle brush, rinse with water and lightly brush. Treat the surface with the final solution, and when the first coating of silver-copper alloy is deposited brush well to obtain a clean metallic surface. A second coating of the alloy may be applied and similarly polished. Apply a coating of shellac to the dried coated surface and then cover with paint.

Galena Blue Mirror (Non-Glaring) U. S. Patent 1.988.663

2 oz.

Solution No. 1

32 oz. No. 2 Potassium Hydroxide, Sodium Hydroxide or Other Similar Alkali Agent 4 oz. Distilled Water 32 oz.

No. 3

Lead Nitrate

Distilled Water

Thiourea (Thiocarbamide) Distilled Water 2 02. 48 oz.

In preparing the above solutions care must be taken to insure complete dissolution of the chemicals and each solution should be shaked well before using In order to produce a lead sulphide film or layer upon the glass or other surface to be treated either of two processes may be employed, one being designated as the "hot" process and the other as the "cold" process.

In either process, the glass or other surface to be coated is initially block

polished or hand rubbed with rouge, after which it is well brushed with water. Following this water brushing operation, a weak solution of tin chloride is applied to the surface to be treated preferably by means of a felt block or bristle brush. The surface is then rinsed well with water and lightly brushed.

The glass so treated is then placed in a horizontal plane and accurately leveled with wedges, the surface to be coated being uppermost. In the "hot" process, after the glass has been initially treated, washed and leveled as just described, the following Solution No. 4 is poured upon the surface to be coated:

No. 4

Distilled Water	4 oz.
Solution No. 1	1 oz.
Solution No. 2	1 oz.
Solution No. 3	1 oz.

Attention is here directed to the fact that in preparing Solution No. 4, the numbered solutions are added to the distilled water in the order given above and that Solution No. 3 is not added until just before the final solution is to be poured upon the glass. Following the application of the tin chloride solution the surface to be treated must be kept wet until the final solution has been applied thereto. As much of the final Solution No. 4 is poured upon the leveled surface as the latter will hold without the solution running over the edges. Heat is uniformly applied to the glass preferably by placing the glass upon a table or bed the surface of which is heated to the required temperature.

In a relatively short time (about 15 minutes) lead sulphide will have deposited out of the final solution and upon the glass. The excess solution is then removed from the glass surface, preferably with a piece of chamois, after which the deposited film is well wiped to obtain a clean metallic surface. A second application of the final Solution No. 4 is then made. In about 10 minutes a second conting or film of lead sulphide will have deposited out of solution upon the first coating, the second coating being also wiped and dried with the chamois. When deposited film of metal shows no dark spots indicating the presence of moisture, a coating of shellac is applied followed by a coating of paint, if desired.

Lead sulphide or galena is a strong metal and adheres tenaciously to the glass. If the mirror shows a grayish color it is usually due to an insufficiently heavy coating of the deposited metal.

An additional coating will remove this defect.

In carrying out the "cold" process, the application of heat is of course omitted and in preparing the final solution no additional distilled water is employed. In other words, the final solution for use in the "cold" process is prepared as follows:

This final "cold" solution is prepared by adding one part of Solution No. 2 to one part of Solution No. 1. These are thoroughly mixed and allowed to stand for about 15 minutes, after which one part of Solution No. 3 is added. After Solution No. 3 has been added, it is necessary to immediately pour the final solution upon the glass due to the fact that the metal tends to deposit out of solution quite rapidly.

Both the hot and cold processes as hereinbefore described have been found quite effective in the application of a firm and homogeneous film or coating of metallic lead sulphide upon a glass surface or the like, it being of course understood that this lead sulphide is formed by the combination of the sulphide present in Solution No. 3 with the lead present in Solution No. 1. It will be understood that in both the hot and cold processes the thickness of the deposited film or coating may be reduced as desired by introducing additional quantities of distilled water either to the final solution or to the primary solutions.

It is important to note that while galena blue (lead sulphide) will not work or combine with silver it will combine with gold.

Aluminum Mirrors

British Patent 433,484

A highly polished aluminum sheet is treated anodically in 24½% borofluoric acid using 20 amp. per sq. ft. at 31 to 33° C., washed and then anodically oxidized in 7% sulphuric acid at 25-26° C. using 12 amp. per sq. ft. After drying, buff with polishing cream.

	Silv	ering	Mirrors		
a. Bi	lver Nit	rate		6	g.
	ater			75	cc.
A	mmonia	(28%))	suffici	ent
T	1 11				

Dissolve silver nitrate in water and add sufficient animonia water to dissolve the precipitate initially formed.

b. Glucose 10 g. Water 100 cc. Mix equal parts of a and b and heat slowly on a steam bath (or in hot water) in the vessel or on the object to be mirrored.

Colored Mirrors

One may use one of two processes to obtain a colored reflecting surface. One process consists of deposition of gold in various thicknesses. The resultant effect of this process is a gold or yellowish to brown colored mirror. This process is limited to a very narrow range of these colors.

A more satisfactory and more widely used process is one where colored glass as used. Pink, red, yellow, purple, green or any desired shade or color glass is used on which silver is precipitated by the regular silvering precipitation process. The silver is then backed on in a normal manner. The resultant effect is a very beautifully colored mirror which is as permanent as the silvering itself. The glass generally used for this purpose is imported.

Of course, one could use a modification of this colored glass process by spraying or brushing on to the front surface of clear glass a colored transparent coating made up of gum sandarae or similar resin in alcohol and dyed to the proper shade. The back of the glass is then silvered in the normal orthodox method. This type of colored mirror is limited in its life by the durability of the front finish coat. It is also very difficult to obtain a uniform smooth reflecting surface by painting or spraying a finish for during the drying period an orange peel effect may manifest itself on the surface and a wavy condition result.

Matte Silver Finish on Watch Dials Formula No. 1

First clean the article well of oil, grease, etc. Then dip into the following solution:

Sodium Dichromate 4 oz. Concentrated Sulphuric Acid 12 oz. Water 1 gal.

The time of dipping depends on the appearance ultimately desired and must therefore be determined by experiment. Rinse well in water, and silver plate in following:

Silver Cyanide 3.5 oz. troy
Sodium Cyanide 4 oz. avoir.
Sodium Carbonate
at least 6 oz. avoir.

Water 1 gal.
Finally soak in boiling water to give dead white color.

No. 2

Precipitated silver is used on some types of high grade watch dials where a dead white matte finish is desired. A raised grain effect is obtained at the same time. The following formula can be employed, using precipitated silver:

Precipitated Silver 1 oz.
Cream of Tartar 2 oz.
Sodium Chloride 2 oz.

Mix dry, add enough water to make thick paste. Apply by running with stiff brush. The proportions may be varied depending upon grain and matte desired. The best results ure obtained on alloys rich in copper such as gilding metal.

Sulphur Resisting Alloy German Patent 591,641

Nickel 44 to 79 lb.
Chromium 9 to 31 lb.
Aluminum at least 9 lb.
Sihcon at least 2 lb.

And 0-14% of one or more of the following: Iron, Molybdenum, Copper, Manganese, Carbon.

Alloys for "Tin" Buttons

Lead	16 g.
Antimony	16 g.
Tin	8 g.

Electrical Resistance Wire Alloy U. S. Patent 1.926.213

Gold 58.4 oz. Nickel 41.6 oz.

Heat Treatment of Aluminum Magnesium Silicon Alloy

Anneal for 1 to 3 hours at 500-550° C.; quench in oil or water and temper at 180-250° C. for 1½ to 3 hours.

Corrosion and Heat Resisting Alloy 35% nickel, 15% chromium (balance iron). This material shows very good resistance to oxidation and corrosion at temperatures up to 2000° F., and still retains an appreciable amount of strength.

Improving Babbitt Metal

Babbitt flow characteristics are greatly improved by adding a small amount of rosin to the molten mass.

Zinc Die Casting Alloys

The following zinc die casting alloys are characterized by low metal cost, ease of casting, excellent finish, good resistance to corrosion, permunence of dimensions, and high strength. The percentage limits apply to die castings. Ingot specifications should be narrower.

U. S. Patent 1,596,761

Zamak-2—A.S.T.M. Alloy XXI—S.A.E. Alloy 921

(The name Zamak is trade marked.)
Aluminum 3.5 -4.5%

| Copper | 2.5 - 3.5% | Magnesium | 0.02-0.1% | Iron | 0.1 % maximum | Cadmium | 0.005% maximum | 1.004 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005 | 0.

Lead 0.007% maximum Tin 0.005% maximum Zinc (Special High

Grade -- 99.99% Pure)

Pure) remainder

This alloy is outstanding in hardness, tensile strength, and resistance to corrosion under severe atmospheric exposure conditions.

U. S. Patent 1,779,525

Zamak-3—A.S.T.M. Alloy XXIII— S.A.E. Alloy 903

Aluminum 3.5 -4.3 %
Copper 0.1 % maximum
Magnesium 0.03-0.08%
Iron 0.1 % maximum
Lead 0.007% maximum
Cadmium 0.005% maximum
Tin 0.005% maximum
Zinc (Special High

Grade - 99.99%

Pure) remainder

This alloy is distinguished by excellent retention of impact strength and dimensions.

U. S. Patent 1,852,441

Zamak-5

Aluminum 3.5 - 4.5 %
Copper 0.75 - 1.25%
Magnesium 0.02 - 0.08%
Iron 0.1 % maximum
Lead 0.007 % maximum
Cadmium 0.005 % maximum
Tin 0.0015% maximum
Zinc (Special High
Grade — 99.99%

Pure) remainder

The characteristics of this alloy are excellent resistance to corresion combined with nearly as high strength as Zamak-2 and retention of dimensions nearly equal to Zamak-3.

U. S. Patent Re 18,600

Zamak-6

| Aluminum | 3.5-4.5% | Copper | 1.0-1.5% | Magnesium | 0.01 % maximum | Iron | 0.1 % maximum | Cadmium | 0.005% maximum | Tin | 0.005% maximum | 0.005% maximum | Tin | 0.005% | Tin |

Zinc (Special High Grade - 99.99%

Pure) remainder

This alloy offers maximum ease of casting at the expense of maximum resistance to intercrystalline oxidation.

Zinc Slush Casting Alloys

The zinc slush casting alloys offer a desirable combination of high strength, good casting finish, case of application of plated and other finishes with low metal cost.

Formula No. 1

Zinc (Special High Grade-99.99% Pure).

This metal offers case of casting and good permanence but lower strength than Formulas No. 2 and No. 3.

No. 2

Aluminum 5-6% Zine (Special High Grade—99.99% Pure) remainder

This alloy offers the greatest case of casting and high initial strength but poor permanence.

No. 3

U. S. Patent Re 18,600

Aluminum 4.55-4.95% Copper 0.65-0.85% Zine (Special High Grade

—99.99% Pure) remainder

This alloy is somewhat hard to cast but has good retention of physical properties and high strength.

No. 4

U. S. Patent 1,596,761

 Aluminum
 5.5
 -6.5%

 Copper
 2.5
 -3.5%

 Magnesium
 0.02-0.1%

 Iron
 0.1
 % maximum

 Lead
 0.007%
 maximum

 Cadmium
 0.005%
 maximum

 Tin
 0.005%
 maximum

Zinc (Special High Grade — 99.99%

Pure) remainder

This alloy offers the highest strength

and permanence of the zinc base slush casting alloys but is also the most difficult to cast.

Zinc Alloy Solders Formula No. 1

Cadmium	82.5%
Zinc	17.5%
Melting Point	508° F.
This solder is mos	t advanta manual

used in soldering zine alloy castings containing aluminum. No flux is necessary.

Note: In making this solder, solid cadmium should be added to molten sine since cadmium fumes have a very dangerous toxic effect. If the cadmium be melted separately, the temperature should not be allowed to rise above 660-700° F. and the surface of the molten metal should be treated with a flux of advantageously | ammonium chloride.

U. S. Patent 1,988,010

		Percentage	by Weight	
Composition	Tin	Zinc	Cadmium	Freezing Point . F.
Formula No. 1	20	53	27	617
No. 2	20	48	32	604
No. 3	30	53	17	630
No. 4	30	46	24	599
No. 5	30	42	28	595
No. 6	40	36	24	581

The above solders are used principally for soldering aluminum and aluminum base alloys. They may be used with or without fluxes depending on the clean-liness of the metal parts.

Cleaning of Zinc and Zinc Alloys

The successful application of plated and other coatings to zinc, zinc alloy die castings, and zinc alloy slush castings depends largely on the suitability and effectiveness of the method of cleaning used.

Cleaning may be accomplished by any one of three methods: (1) Mechanical cleaning by means of sandblasting or scratch brushing, (2) alkaline cleaning and (3) solvent cleaning.

Mechanical Cleaning

Sandblasting with 80 to 100 mesh abrasive is probably most effective since it simultaneously removes grease and dirt and roughens the surface of the metal.

Alkaline Cleaning

Alkaline cleaning has been accomplished very effectively by the use of trisodium phosphate in concentration of 6 oz. per gal. of water. This solution when used at or near the boiling tem-perature and with sufficient current from a 6-volt source to cause violent gassing with the work as the cathode, should remove all grease and oil in ½ to 2 minutes. Alternate hot and cold rines followed by a brief immersion in 10%

hydrochloric acid and a final rinse in hot water to facilitate drying will effectively remove the film of alkaline elemning salts and present a surface suitable for plating or other finishes.

Soldering Zine and Zine Alloy Castings

Zine may be soldered easily, using ordinary solder and a flux consisting of acidulated zinc chloride or killed muriatic (hydrochlorie) acid.

Zine alloys containing aluminum are quite difficult to solder, requiring the uso of a solder consisting of the cadmiumzine entectic (82.5% cadmium-17.5% zinc-melting point 508° F.).

Machining Zinc and Zinc Alloy Castings

Both rolled zinc and zinc alloy castings are machined most advantageously by using tools with more rake than is customary in machining other common metals. The cutting tool should have 15-20° rake and 6-8° clearance.

Two-fluted drills with spiral angles about twice the usual 24 degrees are satisfactory. The included angle of the cutting edges may be advantageously reduced. The clearance angle should be 15 degrees at the periphery of the drill and gradually increased still further as the drill point is approached. Beveling off the end of the flute back of each cutting edge provides more chip clearance for rapid work.

Soapy water is ordinarily a satisfactory lubricant. Kerosene may be used as a lubricant to insure satisfactory separation of chips,

Temperature Glaze for Art Ware and Enameled Brick

Will themselve miles	_
White Lead	35 lb.
Feldspar	17 lb.
Flint	20 lb.
Whiting	8 lb.
China Člay	8 lb.
Colemanite	12 lb.
Tin Oxide	5 lb.
Matte Glaze—Cone 06 to	Cone 02:
White Lead	490 lb,
Whiting	138 lb.
Cornwall Stone	114 lb.
China Clay	210 lb.
Feldspar	98 lb.
Flint	60 lb.
For light groon use 2 to	30% conne

For light green use 2 to 3% copper oxide; for light brown 2% manganese dioxide; for blue 1% cobalt oxide; for vellow 2% sodium uranate; for yellow brown 1/2 to 2% Crocus Martis.

Atware Satin Glaze-Cone 04:

W	hite Lead	410 lb.
Fl	int	227 lb.
Fe	eldspar	85 lb.
	ne Oxide	90 lb.
Ti	n Oxide	60 lb.
B	rium Carbonate	42 lb.
Ti	tanium Dioxide	32 lb.
	ina Clay	54 lb.
	een Matte Glaze-Cone 2	: :
Re	ed Lead	165 lb.
	ldspar	222 lb.
	hiting	40 lb.
	nc Oxide	32 lb.
Co	opper Oxide	12 lb.
	deined Georgia Kaolin	55 lb.
E	nglish Ball Clay	64 lb.
	nis gives a good wax-l	ike textui

Vitreous Enameling Process British Patent 411,380

green for artware or enameled brick.

A mixture of spinel-forming materials, e.g., water 100, ferric oxide 5, nickel oxide 4, calcium fluoride 20, boric acid 45, clay 10 parts, is applied to the iron surface (not necessarily free from rust) and heated at 750-800° for a few minutes in an atmosphere of reduced oxygen content (admixture of producer or waste gases, etc.).

White Vitreous Enamel U. S. Patent 1,933,437

A white enamel for sheet iron and hollow-ware comprises flint 29.236, borax 18.127, sodium nitrate 5.727, sodium carbonate 10.740, red lead 14.920, barium

carbonate 7.757, calcium fluoride 6.563, antimony oxide 4.773, and sodium antimonate 7.160%.

Spark Plugs French Patent 772,601

A ceramic product for spark plugs is composed of a difficultly fusible oxide, e.g., corundum, and a binder which during thermal expansion behaves elastically toward the oxide used. The binder should become plastic at 500-800° C. An example of a binder for use with corundum contains steatite or tale 32.7, kaolin 43.3 and feldspar 24 parts by weight.

Synthetic Precious Stones (Spinels) U. S. Patent 1.952.255

(a) Artificial alexandrite is made by fusing aluminum oxide 85 and magnesium oxide 15% containing colait 0.06, iron 0.04%, and vanadium 0.04%, and (b) a violet spinel by fusing the same aluminum oxide magnesium oxide mixture with iron 1.5 and cobalt 0.005%.

Corundum Abrasive Crystals U. S. Patent 1,966,406

A mixture of raw materials is prepared consisting of aluminous ore such as bauxite or diaspore, silica sand, and an addition agent such as magnesia so proportioned as to give the following ratio

important ingredients:		
Alumina	70	lb.
Silica	25	lb.
Magnesia	5	lb.

This mixture may be fused in an electric furnace of the steel shell are type commonly used in the artificial abrasive industry. The ratio of power input to application of the mix is observed closely as means of governing the temperature of the melt. Thus, under any given rate of power input, a fast feed produces a relatively cool melt, whereas a retarded feed tends to produce a relatively hot bath. The temperature of the melt at the time of withdrawal of the power determines the size and distribution of the corundum crystals. The cool melt pro-duces small crystals uniformly spread through the matrix whereas the hot melt gives rise to the development of large crystals, in pocket formation in the mass.

After the shell has been charged to its capacity and the fusion is completed the electrodes are withdrawn and the cooling process allowed to proceed normally.

MATE	RIALS OF	CONSTRUCTION	239
Brick Glazing		Stone Waterpro	u.fin.a
White Enamel Batch Wei	ghts	An economical treatmen	onng
Red Lead	125.4	durable may be made by	licaciones from
Whiting	35	6 to 12 oz. of a high mel	ting want par-
No. 419 Feldspar	66.1	affin to the gallon of solver	t. such as min-
Raw Kaolin	12.9	eral spirits, naphtha, gaso	line, etc. This
Calcined Kaolin	6.7	usually gives high waters	proofing values
Flint	40	on materials of medium	to coarse tex-
Tin Oxide	30	tures. For fine-pore struc	tures it will be
		desirable to add from 3 to	6 oz. of china
Black Enamel		wood oil to the gallon of	gasoline.
To the above base enamel ba			-
out the tin oxide, the following	is added:		
Cobalt Oxide (CoO)	6	Stucco Waterpro	ofing
Iron Oxide (Fe ₂ O ₃)	8	U. S. Patent 1,9-	12,601
Manganese Dioxide (MnO ₂)	2	Sodium Stearate	5 lb.
		Water	95 lb.
Blue Enamel		Warm to 50° C. and sti	r till uniform.
Batch weights of base enamel:		then add	
Buckingham Spar	66.32	Suct	2 lb.
Red Lead	120.84	Cresol Emulsion	1/4 OZ.
Whiting	36	!	•-
Tin Oxide	57.77		-
Raw Clay (Kaolin)	12.9		_
French Flint	$\frac{40.34}{11.22}$	Masonry Waterpr	
Calcined Kaolin	11.22	British Patent 41	13,463
To the above base is added:		Spermaceti	4 lb.
Black Oxide of Copper	12	Paraffin Wax	1 lb.
Black Oxide of Cobalt Black Oxide of Nickel	18 6	Rubber	1 lb.
Diack Oxide of Nickel	U	Mineral Spirits Trickloroethylens	2550 lb. 25-50 lb.
		Stir until dissolved.	20-00 10.
Brown Enamel		Stir until dissolved.	
To the above base is added:			-
Red Oxide of Iron (Fe ₂ O ₃)) 16	Vitreous Slips for Brick, T	erra Cotta and
The production of other	colors is	Roofing Tile	
merely a matter of experiment	with the		
addition of coloring oxides.	do oo on	Buff	100 11
These glazes contain tin oxicopacifier and on a smooth body		Fireclay Shalo	130 lb. 100 lb.
glossy enamel of sufficient weigh		White Lead	40 lb.
fectly mask the red of the shall		Blue	10 1111
Slips		Ball Clay	200 lb.
	lan mhita	Cobalt Óxide	9 lb.
95% Tennessee Ball Clay (fall slip use English Ball Clay).	or winte	Manganese Dioxide	6 lb.
5% Sodium Chloride are o	f simple	White Lead	50 lb.
materials and easily made up.		Green Ball Clay	200 lb.
For green slip add 20% C Oxide (Cr ₂ O ₃) to the above ba	hromium	White Lead	50 lb.
Oxide (Cr2O3) to the above be	se. The	Chrome Oxide	40 lb.
batch then is:		Manganese Dioxide	24 lb,
Cone 02 to Cone 2		Cobalt Oxide	5 lb.
Clay	380	Black	
Sodium Chloride	20	Ball Clay	60 lb.
Chromium Oxide	80	Blackbird Clay	140 lb.
For blue slip, add 6% Cobalt	Oxide to	White Lead	30 lb.
base. Batch:		Mix the above materials	with sufficient
Clay	380	water to make a heavy all either by spraying or bru	ip, and apply
Sodium Chloride Cobalt Oxide	20 24	dry body, then fire.	usuing on the
ODBIE OXIGE	67	dry tody, then nie.	•

CERAMIC RAW MATERIALS

Chemical Constants

Per Cent Smelt Loss	34.6 22.3 22.3 22.3 34.4 44. 14. 14. 14. 17.3.3 35.5 35.5 17.3.3
Melting Point Deg. C.	Total Part Part Part Part Part Part Part Part
Equivalent Weight	1156 1974 486 1977 486 1977 486 1986 686 1986 686 1987 686 1988 687 688 1988 687 688 1989 688
alent W	BO ₂ BO ₂ BO ₂ BO ₂ BO ₂
	H.O. H.O. S.
Molecular Weight	144.4 118.4
Formula	Al. (OH) Al.
Material	Almminum Hydroxide Almminum Sulphate Antimory Oxide Barium Carbonate Barium Carbonate Black N'ecelle Antimory Bone Ash B

$\begin{array}{c} 59.4 \\ 18.4 \\ 18.4 \\ 18.4 \\ 18.1 \\ 83.8 \\ 16.3 \\ 16.3 \\ 16.3 \\ 16.3 \\ 16.3 \\ 16.1 \\ 12.1 \\ 23.4 \\ 23$	
0.250 0.250 0.250 0.250 0.260	2550
RO ₂ 147.1 186.2 187.2 187.1 187.1 187.1 187.1 187.1 187.1 187.1 187.1 187.1 187.1 188.2 187.2	183.3
RO 294.2 RO 294.2 RO 294.2 RO	
223 2 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	183.3
Pbo Li2C3 Mn02, Mn02, Mn02, Mn02, Mn02, Mn03, Mn	Zr8iO4
Litharge Lithar Garbonate Manganee Dioxide Manganou Chloride Manganou Chloride Magnesium Carbonate (Magnesite) Magnesium Oxide Nickel Sulphate Nickel Sulphate Pottassium Carbonate Pottassium Dichromate Pottassium Nitrate Pottassium Nitrate Releaum Belleau Releaum Riffica (Flint) Sodium Carbonate (Fused) Sodium Carbonate (Carpata) Sodium Nitrate Sodium Nitrate Sodium Nitrate Sodium Nitrate Sodium Nitrate Sodium Nitrate Sodium Sulphate Sodium Sulphate Sodium Sulphate Redium Sulfact Tan Oxide (Stanate) Tin Oxide (Stanate)	Lirconium Silicate

Courtsey of Bureks Plint and Spar Co., Inc.

FUSING TEMPERATURES OF CERAMIC BAW MATERIALS

		Temperature
M aterial	Formula.	Deg. C.
Aluminum Oxide (Alumina)	Al_2O_3	2050
Antimony Oxide	Sb ₂ O ₃	1550
Arsenic Oxide	As ₂ O ₅	200
Barium Oxide	BaO	O ₂ 450
Bone Ash	4Ca ₃ (PO ₄) ₂ ·CaCO ₃	
Borax (Melted)	Na ₂ B ₄ O ₇	732
Boric Acid	B_2O_3	577
Boric Oxide	B_2O_3	577
Calcium Fluoride (Fluorspar)	CaF ₂	1300
Calcium Oxide (Lime)	CaO	257 0
Calcium Phosphate	$Ca_3(PO_4)_2$	1550
Calcium Silicate (Wollastonite)	CaSiO ₃	1540
Cerium Oxide	CeO_2	1950
Chromium Oxide	Cr ₂ O ₃	196
Cobaltous Oxide	C ₀ O	
Copper Oxide (Cupric)	CuO	1235
Feldspar, Potassium	K ₂ O·Al ₂ O ₃ ·6SiO ₂	1170-1235
Feldspar, Sodium	$Na_2O \cdot Al_2O_3 \cdot 6SiO_2$	1120-1215
Fluorspar	CaF ₂	1300
Iron Oxide (Ferric Oxide)	$\text{Fe}_2 \tilde{O}_3$	1565
Kryolith	Na ₃ AlF ₆	
Lead Oxide	PbO	888
Lead Silicate	PbO·SiO ₂	766
Lithium Oxide	Li ₂ O	
Magnesium Oxide	MgO	2800
Manganese Silicate	MnSiO ₃	1273
Manganous Oxide	MnO NiO	1650
Phosphoric Oxide		O ₂ 400
Potassium Oxide	P_2O_5	563
Potassium Silicate	K ₂ O K ₂ O-SiO ₂	red heat 976
Silica (Flint)	SiO ₂	1710
Soda Ash (Sodium Carbonate)	Na ₂ CO ₃	851
Sodium Antimonate	2NaSbO ₃ ·7H ₂ O	931
Sodium Oxide	Na ₂ O	red heat
Sodium Silicate	Na ₂ SiO ₃	1080
Sodium Silico Fluoride	Na ₂ SiF ₆	1000
Tin Oxide (Stannic)	SnO ₂	1127
Titanium Oxide	TiO ₂	1560
Zine Oxide	ZnO	1800
Zirconium Oxide	ZrO ₂	2700

Cold Tile and Brick Glaze U. S. Patent 2,019,980

Portland Cement 10 parts by vol.
1 part by vol. Iron Oxide Calcium Stearate and Water (1-2%) 5 parts by vol.

Mix thoroughly and pass through a screen to remove lumps.

The glaze is now ready for application to the product or article, which, for example, may be cement tile, building blocks or other suitable materials. This blocks, or other suitable materials. This may be accomplished by brushing, dipping or spraying the glaze thereon until the desired coating is effected. The Courtsey of Eureka Flint and Spar Co., Inc.

glazed objects may be trimmed and then glazed objects may be trimmed and then placed in a curing chamber which is kept moist for several days. In order to get best results, the tiles are thereafter placed in storage for a week or longer, to age or cure, until the permanent hard-ening or setting of the glaze is completed.

Enamel Ware Undercoat U. S. Patent 1.962.617

The base metal is sprayed with a suspension of

Cobalt Oxide	3	OZ.
Bentonite	1.5	
Water	100	oz.

CUBICAL COEFFICIENTS OF EXPANSION OF CERAMIC RAW MATERIALS

Material	Formula.		X 10-7
Aluminum Oxide (Alumina)	Al ₂ O ₂	(0.52)	5.0
Antimony Oxide	$\mathrm{Sb}_2\mathrm{O}_3$	(0.02)	3.6
Argenic Uxide	As ₂ O ₅		2.0
Barium Oxide	BaO	(5.0)	
Bone Ash	4Ca ₃ (PO ₄) ₂ ·CaCO ₃	(5.3)	3.0
Borax (Melted)	Na ₂ B ₄ O ₇		3.16
Boric Acid	B ₂ O ₃	(100)	0.1
Calcium Fluoride (Fluoranar)	CaF ₂	(-1.98)	2.5
Calcium Oxide (Lime)	CaO		5.0
Calcium Phosphate	Ca ₃ (PO ₄) ₂		3.6 5
Cerium Oxide	CeO ₂		4.2
Chromium Oxide	Cr ₂ O ₃		5.1
Cobaltous Oxide	CoO 3		4.4
Copper Oxide (Cupric)	CuO		2.2
Fluorspar	CaF,		2.5
Iron Oxide (Ferric Oxide)	Fe ₂ O ₃		4.0
Kryolith	Na ₃ AlF ₆		7.4
Lead Oxide	PhO	(3.0)	4.2
Lithium Oxide	LiaO	(0.0)	2.0
Magnesium Oxide	Mg()	(1.35)	0.1
Manganous Oxide	MnO	(1.00)	2.2
Nickelous Oxide	NiO		4.0
Phosphoric Oxide	P_2O_5		2.0
Potassium Oxide	K ₂ O"	(11.7)	8.5
Silica (Flint)	8(0)	(0.15)	0.8
Sodium Antimonate	2NaSbO ₃ 7H ₂ O	(0.10)	
Sodium Oxide	Na ₉ ()	(12.96)	10.0
Sodium Silicate	Na ₂ SiO ₃	(22.23)	2.96
Sodium Silico Fluoride	Na SiFa		5.0
Tin Oxide (Stannic)	SnO.		2.0
Titanium Oxide	TiO,		4.1
Zinc Oxide	ZnO		1.8
Zirconium Oxide	ZrO _o		9.1

Courtsey of Eureka Flint and Spar Co., Inc.

Pottery Glaze French Patent 44,786

A TOMOR A GUORU	11,100		
Feldspar		26	kg.
Quartz		2	kg.
Minium		49	kg.
Barium Borosilicate		15	kg.

This is applied with coloring materials after grinding, without fritting.

Flooring Tile Norwegian Patent 55,221

The mass before drying is composed of linseed oil 7, coal tar 1, alkalı sılıcate 1, varnish 1, water 1, glue 1, cement 1, quartz sand 5, clay 1 and salt 1 part. all

by weight.

Colored Roofing Granules U. S. Patent 1,944,294

Burned clay granules are impregnated with arsenic trioxide and surface washed. Then treat with 15% basic copper acetate solution, wash and dry. Manufacture of Light-Weight Ceramic Tile

U. S. Patent 1,925,985

A mixture of ball clay 45-65 (56.7), plaster of Paris 10-20 (13.1), and sawdust (1) 25-40 (32.1) is rendered plastic by addition of 80-120 (103)% of water and cast into waxed molds. The dried tiles are heated for 4 hours at about 500° F. until (1) is charred, then slowly (4 hours) up to 1200°, at which temperature they are kept for 4 hours until the carbon is burnt out and shrinkage ccases.

White Enamel for Wire

U. S. Patent 1,938,691

Quench and grind with 8% titanium dioxide.

Light Weight Refractory	Synthetic Lumber
U. S. Patent 1,945,232	U. S. Patent 1,974,277
Brick or Pottery Clay 1 lb. Rice Hull Ashes 2 lb. When the above is fired the product is 40% lighter than usual.	Magnesium Oxide 30 lb. Aluminum Oxide 20 lb. Sawdust 50 lb. Beach Sand 10 lb.

(Continued on page 245;

TEMPERATURE EQUIVALENT OR FUSING POINTS OF CONES

(Adopted from Table XIII, of U. S. Bureau of Standards' Report)

Note: These approximate values are given by the Bureau of Standards to the nearest 5° C. from the average determinations.

Soft Series:

Soft Series:					
	When F	ired Slowly	When Fired Rapidly		
Cone Number	20° C.	per Hour	150° C. p	er Hour	
	° Cent.	° Fahr.	* Cent.	° Fahr.	
022	585	1085			
021	595	1103	605	1121	
020	625	1157	615	1139	
019	630	1166	650	1202	
018	670		660	1220	
017	720	1238	720	1328	
016	735	1328	770	1418	
015	770	1355	795	1463	
014	795	1418	805	1481	
1111		1463	830	1526	
	825	1517	860	1580	
177	840	1544	875	1607	
011	875	1607	905	1661	
Low Temperature Series:					
010	890	1634	00=		
09	930	1706	895	1643	
08	945		930	1706	
07	975	1733	950	1742	
06	1005	1787	990	1814	
0.5		1841	1015	1859	
04	1030	1886	1040	1904	
00	1050	1922	1060	1940	
44	1080	1976	1115	2039	
02	1095	2003	1125	2057	
01	1110	2030	1145	2093	
Intermediate Temperature Series:					
1	1125	2057	7100		
2	1135	2075	1160	2120	
3	1145	2073	1165	2129	
4	1165		1170	2138	
		2129	1190	2174	
	1180	2156	1205	2201	
	1190	2174	1230	2246	
	1210	2210	1250	2282	
	1225	2237	1260	2300	
	1250	2282	1285	2345	
10	1260	2300	1305	2381	
11	1285	2345	1325	2417	
12	1310	2390	1335	2435	
13	1350	2462	1350	2462	
14	1390	2534	1400	2552	
15	1410	2570	1435	2615	
16	1450	2642	1465	2669	
17	1465	2669	1475	2687	
18	1485	2705	1490	2714	
19	1515	2759	1520	2768	
20	1520	2768	1530	2786	

£

TEMPERATURE EQUIVALENT OR FUSING POINTS OF CONES—Continued High Temperature Series:

	Wh	ien Heated at	: 100° per Hou
		° Cent.	° Fahr.
23		1580	2876
26		1595	2903
27		1605	2921
28		1615	2939
29		1640	2984
30		1650	3002
31			3056
32		1700	3092
33			3173
34			3200
35			3245
36			3290
37			3308
38			3335
*39			3389
40			3425
41			3578
42			3659

^{*} The last four cones were heated at 600* per hour.

Moisten with magnesium chloride solution and calcium magnesium chloride and after forming dip in a solution of magnesium silicofluoride and potassium sulphate.

Building Material Austrian Patent 137,323

A fibrous organic material 3, a pulverulent mineral 5-7.5 and water glass solution of 36-38° Bé. 5-7.5 parts are mixed together, molded in a perforated mold, and dried. The organic material may be wood pulp, straw or sugar-cane waste and the mineral may be asbestos or kaolin.

Composition for Floors and Wall Surfaces

Austrian Patent 137,328

Dried sawdust 40-60, cement 30-40 and lime 5-10 parts are kneaded with 50-70 parts of concentrated water glass solution. The dried composition can be subjected to the same mechanical treatments as wood.

Artificial Gypsite Plaster U. S. Patent 1,932,120

Gypsum	26,180 lb.	
Dry Peat	600 lb.	
Clay	2,820 lb.	
O.a.j		

Stir and heat with calcium chloride solution (d. 1.4) 4 qt. in a plaster kettle heating at 155-165° C.

Courtsey of Eureka Flint and Spar Co., Inc. Opal Vitreous Marble, Artificial

French Patent 784,067 Sand 500 kg. Soda Ash 2000 kg. Lume 100 kg. Sodium Nitrate 20 kg.

Artificial Marble Formula No. 1

British Patent 416,774

Diffigu I great area.		
White Cement	50	lb.
Marble Dust	50	lb.
Calcium Carbonate Powder	3.85	lb.
Calcium Oxalate	0.50	lb.
Borax	0.15	lb.
Starch	0.60	lb.
as: at makes and allow	to est	

Mix with water and allow to set.

No. 2 British Patent 430.948

British Patent 450,940	,	
Magnesium Oxide	100	lb.
Marble Dust	30	lb.
Calcium Sulphate Dust		lb.
Make into a paste with m chloride (d. 1.20-1.26) and the	agne	esium Id
Magnesium Oleate Magnesium Stearate Tallow Soap Solution (2%)	1	lb. lb. lb.
TRITOM COURT COURTION (8-70)	**	

STANDARD SCALES FOR TESTING SIEVES

	Di- ameter of Wire (Inches)	25. 25. 25. 25. 25. 25. 25. 25. 25. 25.	0.028 0.033 0.035 0.035 0.023 0.0141 0.0123 0.0100 0.0092
8	Mesh (Per Lineal Inch)		8 8 9 11 11 12 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Tyler Standard Sieve Series Joser ing	Openings (Frac- tions of Inch) (Approx.)	<u></u> - 122524232534533245353451	· 호 호 호 호 호 호 호
Standard	Openings 7 (Milli- meters) (92.25 118.85 113.85 111.33 99.25 6.680 6.680 6.680 6.680 7.560 7.5	2.794 2.794 1.981 1.651 1.1651 1.166 .991 .701 .589 .495 .495 .495 .295
For (Eg G		
Tyler Standard	V2 or 1.414 (Open- ings in Inches)	7.42 .525 .525 .371 .371 .185	.093 .065 .046 .046 .0232 .0232 .0164
	Wire Diameter (Milli: meters)	25	3 4 2 2 3 3 4 2 3 3 4 2 3 3 4 2 3 3 4 2 3 3 4 3 4
	— .		
Series	Wire Diameter (Inches)	600.00.00.00.00.00.00.00.00.00.00.00.00.	0165 01160 01130 01130 01130 00087 00064 00055 00055 00050 00050 00050 00050 00050
d Sieve E	Sieve Opening (Milli- meters)	666 67.5 7.5 7.5 7.5 7.5 7.5 7.5 7.5 7.5 7.5	#81.159 150 150 150 150 170 149 105 105 105 105 105
U. S. Standard Sieve Series	Sieve Opening (Inches)	2655 223 223 223 157 111 0033 00469 00469	.0331 .0238 .0238 .0197 .0165 .0138 .0033 .0049 .0049
Ģ.	Sieve Number		250 250 250 250 250 250 250 250 250 250
	Meshes per Lineal Inch	2288228828232	20.16 27.62 32.15 32.15 38.02 44.44 52.36 61.93 72.46 85.47 100.01 120.48

								_		
.0056	.0042	.0038	0008	000	.0021		807	100	140	
80	100	115	150	170	200	sizing-3 to 11/2 inch opening	1	ı	I	le has as its base an opening of .0029 ing in 200-mesh .0021-inch wire, the stan by the U. S. Burean of Standards, the the ratio of the square root of 2 or 1.41st. sizing is required column 2 shows the challenges in the ratio of the fourth root courters of Eureka Finst and Spar Co.
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.175	.147	.124	104	.088	.074	zing-3- t	i	1	i	ing in 200-mesh .0021-inch wiften 200-mesh .0021-inch wiby the U.S. Burean of Stan Her ratio of the square root of sizing is required column 2 . Scale with intermediate saleves. Increase in the ratio of the Courtery of Eureka Flint.
6900.	8000	.0049	.0041	.0035	.0029	coarser	1	1	1	scale has a pening in 2 sed by the in the ratio 3ser sizing in Scale will has increase
100	0000	1 8	T#00*	18	.0029	For	ı	ı	ı	This sieve scale has as its base an opening of which is the opening in 200-mesh .0021-inch wire, the sieve, as adopted by the U. S. Bureau of Standards, ings increasing in the ratio of the square root of 2 or 1. Where a closer sizing is required column 2 shows Standard Screen Scale with intermediate sieves. In the sieve openings increase in the ratio of the fourth or 1.189.

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.062 .053 .044	P 4 4 7 7 7
.246 .0024 .0021 .0017	essentiall the basic are relativation of the In making be done
.0097 230 270 325	mm. is the series 32 as the ppening. that this selection o
238.10 270.26 323	This sieve opening of 1 below this in 70 f 2, or 1.189 next smaller or recommended example, the sin the series.

Stone Wood Composition Flooring British Pater 426,739

A 1:9 mixture of sodium thiosulphite and calcium carbonate is added to plaster of paris containing sawdust or cork filler. Proportions by volume of 1:5:4 respectively are preferred.

Artificial Stone Flooring U. S. Patent 1,968,784

 "Eternit" Artificial
 Slate

 Cement
 100 lb.

 Aslextos
 20-25 lb.

 Pigment
 5 lb.

 Rosin Solution
 to suit

Artificial Stone

British Patent 430,404

(a) Portland coment 2 and clay dust 025 are mixed, (b) cement 2 and slate dust 1 are mixed therewith, (c) cement 2 and shale 1 are mixed therewith, (d) cement 2 and river of sea sand 1 parts are mixed therewith, (e) the calcium oxide solution is added, 0.6 gal. at a time, with mixing after each addition. (7 lb. of calcium oxide and 2 gal. of water may be used for each 28 lb. of cement), (f) the product is molded into slabs, dried 24 hours and then baked 2 hours at 100° C. The slabs are then painted, heated 1 hour, cooled 30 minutes, coated with enamel or cellulose paint, heated 2 hours at 150°, smoothed with cuttlefish bone and polished.

Wall Board, Artificial U. S. Patent 1,976,190

Calcined magnesite 12 parts; sawdust 3 parts; an aqueous solution of magnesum chloride at about 18° Bé., 14 parts; molasses ¼ of 1 part; said ingredients being combined in a creamy fluid mixture sufficiently thin to be readily poured into a form or mold.

Commercial Porcelain

The clay used in making porcelain varies in each locality and it will thus be necessary in the following formulas to include the chemical analysis of the clay

and spar used. The physical properties of various mixtures are best illustrated by the triangular diagram shown by Gilchrest and Klinefelder in the Electric Journal, March and April, 1918. This diagram shows the variation of mechanical, electrical and thermal properties with variation in mixture.

The usual mixtures for electrical porcelain is 40-50% clay, 25-30% spar and 25% quartz. The European insulator materials are ground extremely fine and fired to a hard glass-like body, usually Seger cone 14-16.

German Porcelain Mixtures Insulator Porcelain

	* ATCOUNTE	
Kaolin from Halle		50.4%
Clay from Halle		31.5%
Spar		18.1%
T1 1		

Equal to 53% clay substance, 29% quartz and 18% spar.

Household Porcelain

Formula	No. 1	No	. 2	No. 3
Kaolin from Halle	55%	60	%	22.4%
Clay from Halle	27%	_		44.8%
Zettlitzer Kaolin	_	18	%	16.2%
This canala				

Clay Substance 54% 55 % 60 % 22.5% 22.5% Quartz 28% 22.5% 17.5% Spar 18%

The kaolin from "Halle" mines contains about 61.77% clay substance, 37.84% quartz and .39% spar. The clay from the same mines are about 70% clay substance and 30% quartz.

Karlsbaden Czechoslovakian Porcelain Clay Substance 52 % Quartz 29.62%-24.5 % Spar 17.26%-21.93% 1.25%- 1.6 %

Calcium Carbonate

Danish Porcelain

Clay Substance	31.8%
Quartz	30.8%
Spar	33 %

Chinese Porcelain

	01 0040444	
Clay Substance Quartz		31.8% 30.8%

.11 Potash German Glaze

.67 Lime Aluminum Oxide + 10 Silica .22 Magnesium Oxide

The above glaze is made from .11 Potash + .11 Aluminum Oxide + .66 Silica = 61.27 lb. Spar .67 Lime

lb. Marble = 67 .22 Magnesium Oxide as Magnesite = 18.48 lb. Magnesite .89 Aluminum Oxide + 1.78 Silica = 230.5 lb. Zettlitzer Clay

7.56 Silica = 453.6 lb. Hohenbacker Sand

Natrium Spar 19.4% Mica 18 % The firing temperature of the German

and Danish porcelain varies between Seger Cone No. 14 and No. 16. The household china is usually bisquit fired at a temperature of about 900° C. before glazing and then glazed and given the final firing at about 1400° C. They are then painted or decorated and given short firing at about 600° to set the colors.

The quartz used chiefly in the above mixtures comes from Sweden, the feldspar from Norway, whereas some of the best clays come from Czechoslovakia, although good raw materials are obtainable in many countries.

The chemical analysis of the above materials is as follows:

Zettlitzer Clay 12.65% Silicic Acid 46.9 % Aluminum Oxide 38.56% Ferric Oxide .84% Potash and Sodium Oxide 1.05% Giving the following technical analysis Clay Substance 98.8 % Quartz and Spar 1.2 % **F**eldspar Nor-Czecho-

wegian slovakian 62.25% 54.5 % Silicic Acid 54.5 % 19.96% Aluminum Oxide 19.75% Ferric Oxide .35% 1.75% Potash (K2O) 14.32% 11.5 %

Lime (CaO) .55% Magnesia (MgO) .21% Sodium Oxide

(Na₂O) 1.36% Magnesia Porcelain

porcelain usually consists of about 85% powdered talcum and 15% settled gelantinic magnesium silicate, or 80% talcum and 20% China clay. These magnesia porcelains have very good electrical and mechanical properties and extremely small shrinkage during firing allowing the pieces to be made to very close dimensions. They also retain their high electrical resistance up to very high temperatures. The chemical formulas for some of the

glazes used by various European manu-facturers are as follows:

Another clear and fine German glaze	
.3 Potash .7 Lime 8.8 Aluminum Oxide + 8 Silica]
Danish Glaze	
.65 Potash .35 Lime Aluminum Oxide + 15 Silica	9
This glaze is made of	. :

China Clay 6.75 lb. Quartz 48.75 lb. Spar lb. Crayon 2.75 lb. Bisquit fired porcelain (Powdered) 13.75 lb.

All the above formulas are based on pure European porcelain materials and if materials obtained locally are used a thorough chemical and rational analysis must be made of the raw materials used and the formulas corrected for the varying compositions of the materials.

Low Expansion Borosilicate Glass U. S. Patent 2,012,552

A borosilicate glass having a thermal coefficient of expansion of about .000005 and consisting essentially of silica 72%, magnesia 12%, boric oxide 8%, sodium oxide 6% and potassium oxide 2%.

Ultra Violet Stable Glass British Patent 424,366

Potassium Carbonate	13.77	lb.
Potassium Nitrate	6.71	lb.
Calcium Carbonate	8.93	lb.
Barium Carbonate	3.22	lb.
Magnesium Carbonate	18.53	lb.
Boron Oxide	31.04	lh.
Aluminum Oxide	28.80	lb.
Diammonium Hydrogen		
Phosphate	48.70	lb.

Brown Glass

U. S. Patent 2,014,230

A batch for making brown glass comprises in addition to the ordinary glass composition 0.5 to 3.0% of ammonium sulphate and 0.5 to 5.0% of organic matter.

Coloring Glass Austrian Patent 140,547

Colored coatings are produced on sulphide glass not containing free carbon. A typical sulphide glass is made from Sand 87 lb.

20 lb. Soda Ash

Potassium Carbonate Lime	10 lb.
Borax	11 lb. 2 lb.
Ferrous Sulphide	3 lb.

Colored Continu Composition

00.00	ica conting compe	DICTON	
Cuprous		30	lb.
Calcined	Copper Sulphato	30	lb.
Calcined	Clay	120	lb.

Luminescent Glass British Patent 415.536

Zine sulphide and/or cadmium sulplude, etc. are/is either added to the class or formed in the glass by reduction the corresponding sulphates with zinc, in, magnesium powders, carbon, sulphur, etc. or by combination of the oxides or carbonates with sulphur. The presence of 0.01-0.4% of a heavy metal (cadmium, copper, antimony, manganese, etc.) is also necessary. An orange-yellow glass is composed of silicon dioxide 66, aluminum oxide 3, borie anhydride 3, calcium oxide 3, zinc oxide 5, potassium oxide 5.5, sodium oxide 11.5, manganese sulphide 0.63, and zinc sulphide 2.37%.

Cream Colored Opaque Glass U. S. Patent 1,956,176

Fuse together	
Sand	885 lb.
Soda Ash	306 lb.
Feldspar	675 lb.
Cryolite	90 lb.
Calemu Fluoride	50 lb.
Sodium Nitrate	30 lb.
Arsenic Trioxido	4-10 lb.
Ferric Oxide	4-10 lb.
Sodium Uranate	2-7 lb.
Belenium	⅓-₩ lb.

Vacuum Tube Glass

U. S. Patent 1.969,277

Boric Oxide	40	to	60	lb.
Sodium Oxide	4	to	5	lb.
Calcium Oxide	10	to	11	lb.
Alumina	11	to	13	lb.
Bilica	20	to	30	lb.

Lightly "Frosted" Glass

Gelatin Bodium Water	Fluoride	4.5 2 30	g. cc.

The gelatin is first dissolved in the water and then the sodium fluoride is added. The solution is then poured over a glass plate and the latter is allowed to dry in a horizontal position. When completely dry, the plate is immersed in a dilute solution of hydrochloric acid for 30 seconds, and is then again allowed to The remainder of the gelatin may then be removed with the aid of hot water.

Acid- and Waterproof Cement U. S. Patent 1,973,731

Silicate cements are rendered harder and denser by the addition of 1/2 to 2% of aluminum or calcium hydroxide and sodium silico-fluoride.

Special Cement French Patent 777,055

Portland Cement Clinker Slag Silica Slaked Lime Plaster Stone	50-55 kg. 8-12 kg. 8-12 kg.
Grind all together.	2-6 kg.

Cellular or Light Weight Concrete U. S. Patent 1.985,905

A slurry is formed by mixing coment with following foam producing compound:

Water	500 lb.
Casein	100 lb.
Slaked Lime	25 lb.
Benzaldehyde	7 lb.
Beta Naphthol	1 lb.
Arsenic Trioxide	1 lb.

Coloring Concrete

For coloring white Portland cement, 5 to 10% of the following materials are generally employed:

Iron Oxides Red, yellow, brown, black Manganese Dioxide Brown, black Chromium Oxide Green Ultramarine Blue Blue Cobalt Blue Blue Carbon Pigments

Black Certain types of pigments such as those containing Prussian blue, zinc and lead chromates, and cadmium lithopone cannot be used. Chrome green needs to be carefully distinguished from chromic oxide green. Lead oxide pigments are unsuitable and ultramarine is not entirely stable. The fading of colored concretes is due to the formation of a film of calcium carbonate on the surface.

Fire Resistant Concrete Hungarian Patent 109,616

3 qt. Chamotte Flour (10 mil.gr.) Cement 1 qt. Quartz Powder 1 qt.

Waterproofing Mortar and Concrete Austrian Patent 138,387

10 kg. Olein Ammonia (0.910) Mix until uniform and then add slowly

with stirring Aluminum Sulphate (22° Bé.) 2 l. or Zinc Oxide

In use, the above mixture is added to 100 times its weight of 20% milk of lime and the latter is used in place of the water to be used with the cement.

Flexible Paving Material U. S. Patent 1,961,678

Approximately 60% of coarse (4.14in.) anthracite bone and rock from a cleaning plant together with fillers, e.g., sand 30 and marble dust 5%, is mixed with 6-12% of a bituminous binder.

Road-Surfacing Material Swedish Patent 80,677

Slabs for road, sidewalk and floor surfacing are made from a mass consisting of 20.4% wood tar, 20.4% coarse sand below 3 mm. size, 40.8% fine sand having a grain size of 0.25-2.0 mm., 4.1% ground unslaked lime, 8.2% cement and 6.1% of fireclay.

Tennis Court and Path Surfacing British Patent 430,001

Twelve pounds rosin are mixed hot with 1 gal. raw linseed oil and 1 oz. powdered alum, 2 gal. of the resulting syrup being mixed with 6 cu. ft. dry sand and 30 oz. chrome green being added. If a quick-setting, tough material is required, 70 oz. of tung oil and 5% (calculated on total oils) of a 4% cobalt linoleate are added.

Asphalt Powder German Patent 613,620

Asphalt (M.P. 45° C.) Glass or Mica Powder 3 lb. Warm and mix, cool and powder.

Pavement Joint Packing U. S. Patent 2,016,404

Rubber	40 lb.
Asphaltum	7 lb.
Whiting	46 lb.
Sulphur	3 lb.
Ammonium Carbonate	2 lb.
Work into a porous mass an	d cure by
heating.	•

Refractory Compound British Patent 413,398

A mixture of refractory plastic clay with finely ground glass (of any quality) borax and sodium chloride (e.g., 20, 2, 1 and 3 parts by weight respectively) yields refractory products of increased durability and is also suitable for use as a refractory plaster or cement.

Ingot Mold Refractory U. S. Patent 1,984,759

	,
Chrome Ore	8-10 lb.
Basic Slag	2- 5 lb.
Magnesite	10-12 lb.
Calcined Fire Clay	50-30 lb.
Plastic Clay	10-15 lb.
Common Fire Clay	20-28 lb.

Spark Plug Refractory British Patent 422,474

Corundum		96	lb.
Titanium Dioxide		2	lb.
Magnesium Dioxide		2	lb.
Heat; grind; mix with	a	little	acid,

mold and fire at 1630° C.

Refractories Resistant to Spalling

Bricks for suspended arches of boiler furnaces can be made of a highly aluminous clay containing silica 54.48, aluminum oxide 43.18, ferric oxide 1.10, calcium oxide 0.86, magnesium oxide 0.18%; ignition loss was 0.32%. No plastic clay was added.

Fused Silica, Improved U. S. Patent 1,984,178

An insulating composition having essentially the properties of fused silicable but being characterized by improved workshility when plastic and decreased brittleness, consists mainly of silica and contains as constituents about 1/4 to 11/4 per cent of beryllium oxide and about 1/4 to 2% of aluminum oxide and about 1/4 to 2% of aluminum oxide

Inorganic Electric Insulation for Steel U. S. Patent 1,951,039

Steel sheets are coated with a mixture

f	
Lime	15 lb.
Iron Oxide	28 lb.
Sodium Silicate	70 lb.
Water	200 lb.
Bake at 240° C. and anne	sal at 800° C.
Bake at 240° C. and anne	sal at 800° C.

Tooth Stump Model for Dental Crowns British Patent 421.872

Dittion I decide 421,012		
Aluminum Oxide	50	OK.
Silica	16	OK.
Calcium Sulphate	33	OZ.
Calcium Sulphate Gold Chloride Solution (1%)	1	0 2.

Insulating Decorative Molding British Patent 430,041

Acid	15	lЬ.
	8	lb.
	20	lb.
	65	lb.
	15	lb.
to make	plas	tic
	Acid	Acid 15 8 20 65

Sound Absorbing Composition U. S. Patent 1,996,032

Mineral Wool	851/4	lb.	
Glue	2	lb.	
Cooked Starch	9	lb.	
l'yrophyllite	21/2	lb.	
Beta Naphthol	1/2	OZ.	
Aluminum Sulphate	2	OZ.	

Treating Peeled Rattan U. S. Patent 1,959,462

The plugs are impregnated with a 1% aqueous solution of glycerol, water is evaporated and the treated plug is sprayed with a solution formed of cellubid 2 lb, and acctone 1 gal. to which powdered aluminum 20 g. and powdered zinc 3 g. have been added, to serve as a scaling and preservative agent.

Minimizing Wood Shrinking and Swelling

Soak wood in water in a vacuum chamber, the air being removed by alternate evacuation and breaking the vacuum. Soak for a week in "Cellosolve" and then distil under vacuum of 60 cm. mercury at 40-45° C. in a number of steps over a period of 3 days. Bgy, distil at 100° C. The "Cellosolve" may be sub-

sequently replaced, if desired, by soaking in oil or molten wax for more than a week at temperatures up to 85-90° C.

Wood Antiseptic and Fireproofing British Patent 425,495

Combined fireproofing and preservative properties are claimed for mixtures in aqueous solution of a metallic phosphate, a borate, and a chloride. Impregnation of wood can be undertaken in the usual metal apparatus, since the ingredients are without chemical action on iron. Being resistant to temperatures up to 1000° C., the materials specified not only prevent spread of combustion, but | it.

smother flames entirely. These preparations are also said to be suitable for preserving and fireproofing paper, fabrics, etc., by the simple process osaking. An example of a water-insol-uble preparation comprises 5 lb. dibasic sodium phosphate, 3 lb. sodium tetraborate, 1 lb. zinc chloride, 12 lb. 25% aqueous ammonia solution, and 90 pt. (maximum) water.

Fireproofing for Wood

Ammonium Phosphate 100 kg. 10 kg. Boric Acid 1000 1. Water

Mix and dissolve and immerse wood in

Hardness Scale

1. Talc 2. Rocksalt 3. Calcite	4. Fluorite 5. Apatite 6. Feldspar	8. Topaz 9. Corundum 10. Diamond
	7. Quartz	

Hardness of Materials

The above numbers give only the order of arrangement as to hardness.

Agate	7.	Hematite	6.
Alabaster	1.7	Hornblende	5.5
Alum	2-2.5	Iridium	6.
Aluminum	2.	Iridosmium	7.
Amber	2-2.5	Iron	4-5.
Andalusite	7.5	Kaolin	1.
Anthracite	2.2	Lead	1.5
Antimony	3.3	Loess (0°)	0.3
Apatite	5.	Magnetite	6.
Aragonite	3.5	Marble	3-4.
Arsenic	3.5	Meerschaum	2-3.
Asbestos	5.	Mica	2.8
Asphalt	1-2.	Opal	4-6.
Augite	6.	Orthoclase	6.
Barite	3,3	Palladium	4.8
Beryl	7.8	Prosphor Bronze	4,
Bell-metal	4.	Platinum	4.3
Bismuth	2.5	Plat-Iridium	6.5
Boric Acid	3.	Pyrite	6.3
Brass	3-4.	Quartz	7.
Calanime	5.	Rock-Salt	2.
Calcite	3.	Ross' Metal	2.5-3.0
Copper	2.5-3.	Silver Chloride	1.3
Corundum	9.	Sulphur	1.5-2.5
Diamond	10.	Stibnite	2.
Dolomite	3.5-4.	Serpentine	3-4.
Feldspar	6.	Silver	2.5-3.
Flint	ř.	Steel	5-8.5
Fluorite	i.	Talc	1.
Galena *	2.5	Tin	1.5
Garnet	7.	Topaz	8.
Glass	4.5-6.5	Tourmaline	7.3
Gold	2.5-3.	Wax (0°)	0.2
Graphite .	0.5-1.	Wood's Metal	3.
Gypsum	1.6-2.	Zinc	2.5
4 E	2.0 4.		4.0

Wood Preservative British Patent 424,941

On impregnating wood with a mixture of a chromate, a salt of a heavy metal—i.e., a metal with a specific gravity greater than 4—and sodium fluoride, a reaction is claimed to take place in contact with the acids and the cellulose in the wood with formation of water-insoluble substances exercising powerful fungicidal action. A preferred mixture comprises 50% potassium or sodium bichromate, 30% zinc chloride, and 20% sodium fluoride, and the impregnation can be effected by standard methods such as a vacuum and pressure process, using a 1% aqueous solution.

Wood Preservative British Patent 425,781

Boric acid and ammonium dihydrogen phosphate may be added for fireproofing.

Creosote	Wood	Preservative	Emulsion
Glue Sulphor	nated Fa	atty Alcohol	0.08 g. 0.02 g.

Creosote
Water

Allow first two items to swell in water
and then mix with creosote and run
through colloid mill. Stability is improved by neutralizing any free acidity

in creosote with alkalı.

Cresylic Wood Impregnation Bath
Cresylic Acid 100 lb.
Red Oil (Double Pressed) 100 lb.
Cunstic Soda Solution 32° Bé. 20 lb.
Manipulation: Add caustic soda solution to red oil at 50° C., add cresylic acid
slowly with constant agitation and cool
rapidly.

Arsenic Cement Coating for Wood Piling
Sand 12 lb.
Cement 3 lb.
Arsenic, White 1 oz.
Mix dry and add water before use.
Then apply to piling by air gun.

Oil for Wood Preservation
Carbolneum, Pale (Bleached
with Chlorine, Tar Oil)
Rosin, Pale
Amilia Dye, Oil-Soluble
Linseed Oil
Drier
Toption Soluble

to suit
Linseed Oil
Drier
Toption Soluble

to suit
Linseed Oil

PAPER

Paper Coating

Formula No. 1

Cool to 35° C. and add Ammonia (28°) Cold Water to make

2 lb.

50 gal.

roman 110. 1	Cold Water to make 50 gal.
Argentine or Silver Paper:	The emulsion should be allowed to
Argentine Pulp 40% 90 lb.	stand for at least 24 hours before use as
Casein Solution	it seems to improve with age. This emul-
(1/4 lb. per gal.) 21/3 gal.	sion is added to the coating mixture in
Carnaula Wax Emulsion 1/2 gal.	sufficient amount to give the desired gloss
Toluol 1 pt.	when the paper is flinted.
Carbon Tetrachloride 1 pt.	
Nigrosin 9 oz.	No. 4
The casein solution is made as follows:	Canadian Patent 344,222
Casein 62 lb.	Phthalic acid (8.5) and caustic soda
Borax 7 lb.	(5.5 parts) are dissolved in 1300 parts
Trisodium Phosphate 7 lb.	of water at room temperature. White
Water to make 50 gal.	molding plaster or calcined gypsum (850
The carnaula wax emulsion is made	parts) is added and the mix is stirred for
with this formula:	1 hour. To this slurry is added 1100
Carnauba Wax 140 lb.	parts of casein glue containing 170 parts
Castile Soap 20 lb.	of dry casein. The product is used di-
Water to make 140 gal.	rectly on the paper-coating machine. The
No. 2	method may be modified for the utiliza-
A coating mixture which will give a	tion of a mixture of the deflocculated
high finish when calendered is made up	gypsum and coating clay by using soda ash as the electrolyte, and Turkey-red oil
as follows:	may be added to the final product.
Water 65 gal.	may be added to the man product.
Soda Ash 3 lb.	
Ammonia 4 gills	Playing Cards
Satin White Pulp 440 lb.	British Patent 405,502
English Clay 650 lb.	The cards are composed of a core of
Stir untill thoroughly mixed and	textile fabric impregnated with a solu-
smooth and add the following casein	tion of cellulose derivative and coated on
eolution:	both sides with a layer or layers of cellu-
Water 50 gal.	lose derivative solution containing such
Casein 100 lb.	a small amount of plasticizing agents
Soda Ash 10 lb.	that the cards are clastic. A suitable
Trisodium Phosphate 7 lb.	composition consists of cellulose acetate
Borax 5 lb.	2.5, acetone 4, denatured alcohol 6, cas-
Ammonia 6 gills	tor oil (plasticizer) 0.35 and dry pigment
This coating mixture will produce a	0.16 kg. To make the card opaque the
high finish when calendered, that is suit-	composition used for coating one side
able for the highest grade lithographic	may contain metallic pigments, e.g.,
or process printing.	bronze powder.
No. 3	S4 2 Cl 4
Wax Emulsion for Flint Paper	Stencil Sheets
Yellow Laundry Soap 7 lb.	U. S. Patent 2,004,484
Carnauba Wax 50 lb.	Yoshino paper is coated with
Water 12½ gal.	Gelatin 13 oz.
Boil with live steam till thoroughly	Hard White Soap 42 oz.
emulsified (from 3-4 hours).	Almond Oil 56 oz.
2	54

Treating Parchment Paper for Wrapping Butter

Parchment for salt butter is immersed for ten minutes in a solution of 2½ lb. salt in 10 gal. water heated to 220° F.

Separating (Non-Sticking) Paper U. S. Patent 2,017,449

A flexible fibrous sheet is coated with Sodium Silicate 140 g. (dlycerin 15 g. Carnauba Wax Emulsion 1 g.

Gummed Paper U. S. Patent 1,940,363

A thin film of adhesive composed of 90% of dextrin and 10% of gelatin glue applied to transparent paper enables it to be printed with common quick-drying inks and to adhere to gluss.

Waterproofing for Paper

Trihydroxyethylamine	-
Stearate	41/4 lb.
Stearic Acid	1/2 lb.
Water	100 lb.
Boil and mix until smoo	th; pour inte

this slowly while stirring vigorously
Paraffin Wax (Heated
to 90-100* C.) 30 lb.

Stir until cool.
Use 1 part of above emulsion to 5-10 parts of warm water.

Non-Staining Waterproofing for Paper U. S. Patent 1,968,907

Petrolatum Wax	2590	
Ester Gum Paraffin Wax	5-75 5-50	

Waterproofing for Paper Australian Patent 5604

Shellac	22 oz.
Alcohol	75 oz.
Formaldehyde	3 oz.

Waterproofing Paper and Fiber Board Canadian Patent 343,302

The strength and water resistance of a liquous material are increased by beating in a liquor containing % to 4 lb. of casein per 100 lb. of pulp lime from 10 to 25% of the weight of the casein, and sodium fluoride from 5 to 12.5% of the weight of the casein. The material treated may be paper, fiber board, as-

bestos board or the like. The strength and water resistance may be increased if a relatively small quantity of formaldehyde is added to the treating solution. If the fiber so treated is somewhat too brittle, a softening agent such as glycerol, sulphonated or saponified oil or fat may be added to the treating composition.

Waterproofing Paper and Textiles U. S. Patent 1,981,405

	-,,		
Glue	15	05.	
Water	83	OZ.	
Formaldehydo	1-2		

Dissolve glue in water and mix formaldehyde with it vigorously and spray immediately on material to be waterproofed.

Embossed Waterproof Wallpaper U. S. Patent 1,936,355

Stearic Acid	4 lb.
Japan Wax	5 lb.
Triphenyl Phosphate	8 lb.
Dibutyl Phthalate	1 lb.
Heat to 90° C, and add	
Water Shellac (40%)	56 lb.
Triethanolamine	2 lb.

Cool to 70° C. and add successively with vigorous stirring

Anmonia (28%) 1 qt.
Water 3 gal.
Anmonia (28%) 3 qt.
Water 3 gal.
Latex + 4% Sulphur 3 lb.
Water to make 28 gal.

Odorless Greaseproof Paper and Textiles British Patent 431,191

This composition comprises a cellulose derivative and rubber or chlorinated rubber dissolved in a solvent free from benzene or its derivatives and containing di, ri, or per-chloroethylene and/or methylene chloride. The composition may be employed for the production of artificial silk, filaments, threads, films, sheets, and the like, in which case the preferred proportions are chlorinated rubber 30 to 50 parts and cellulose derivative (nitrate or asstate) 800 to 900 parts. A typical solvent for such a mixture comprises trichloroethylene or methylene chloride 180 to 300 parts, and acetone 2000 to 3375 parts. A further application of the composition is in the production of an odeless and grease-proof wrapping paper, and of coated textile and like sheets. A

switable composition for this purpose comprises chlorinated rubber 15 to 20 parts, cellulose nitrate or acetate 66 to 80 parts, dissolved in a mixture of trichloroethylene 90 to 120 parts and acetone or methylene chloride 1000 to 1300 parts. To this composition may be added a mixture of diethyl phthalate, castor oil and parafin oil as plasticizer. A paper base may be coated by passing it through the composition, which is maintained at a temperature of 28-38° C. and the conting dried by passing through a drying chamber. The drying step is preferably followed by a humidifying operation by passing the coated paper through a tower containing humidified air. In place of the cellulose acetate or nitrate there may be used benzyl cellulose. The rubber and the cellulose derivatives may be dissolved separately and the solutions mixed.

Wax Size, Paper Formula No. 1 U. S. Patent 2,009,488

First emulsify a corn oil soap with water to form a paste. Next mix into this paste modified starch in the ratio of preferably approximately about 15 parts of modified starch to 10 parts of corn oil soap. Thereafter, and while the mixture of corn oil soap and modified starch is constantly agitated, incorporate a wax, preferably melted parafin, although other waxes such as montan, japan, carnauba, etc., may be used alone or in substitution for a portion of the paraffin. The wax may be incorporated in the amount of 75 parts to 15 parts of modified starch and 10 parts of soap.

The mixture thus produced may be incorporated in the beaters in which event add a small percentage of paper manufacturers' alum to aid in the precipitation as the retention of the size is increased in this way. The mixture thus produced may also be used as a surface sixing and so used as mixed with sufficient water to produce the desired fluidity. The amount of water equal to the weight of the wax component is satisfactory.

The corn oil soap prevents foaming in the compounding of the size and the modified starch eliminates to a large degree the softening effect upon the paper heretofore produced through the use of wax emulsion sizes.

The resulting size paper has a high finished hard surface and the sizing is equally applicable to cellulosic and as-

bestos paper stocks. In connection with asbestos paper, the resulting size renders the paper highly water resistant.

No. 2 Canadian Patent 352,422

Pulp Fiber (Dry Weight) Water	1,000 20,000	lb. lb.
Mix in a beater and add		
Calcium Carbonate	300	lb.
Ammonium Resinate		
(Dry Weight)	15	lb.
Water	500	lb.
Alum	15	lb.

Plant Cover and Fruit Wrapping Paper Canadian Patent 346,222

To each ton of unbleached sulphite pulp is added 160 lb. of thick size, or other suitable size equivalent to 112 lb. of dry size. The stock is beaten for 30 minutes; then 40 lb. of copper sulphate in suitable water solution is added to the stock. Beating is continued for 15-20 minutes. A slight excess of size is maintained with a backwater pH of not less than 6.0. The paper prepared from the stock will contain an excess of the desired 1% per weight of copper resinate; that amount of copper resinate being considered necessary to impart to the paper sufficient resistance to the deterioration and destruction of its fiber when used as a plant cover or fruit wrapper.

Detecting Artificial Watermarks in Paper

Artificial watermarks produced by impression on the nearly dried paper with a rubber stamp are differentiated from the genuine by sprinkling the area with a mixture of 100 g. of dry icing sugar and 0.5 g. of concentrated Rhodamine-6G, placing the paper in a dish of water, and examining in filtered ultra-violent light. The design of genuine watermarks is marked for a few seconds by a bright golden fluorescence, which is absent in the case of artificial watermarks.

Discharge Effects on Tissue Paper

Discharge effects on tissue paper are produced in a very simple manner by passing the tissue paper through the solution of an easily dischargeable dyestuff in the dyeing machine, and spraying or printing on a solution of 1 lb. Hydralite C extra per 1 gal. water to which has been added a solution of 3½ oz. acetate of zinc per 1 gal. water or 1 pint acetate

of alumina of 18°	Tw.;	the	paper	is	then
dried quickly.					

U. S. Patent 1,997,487

An absorbent paper is treated with Glue 6 oz. Formaldehyde 3 fl. oz. Water 10 make 1 gal.

Transfer Printing Paper U. S. Patent 1,965,257

Rubber Latex (60%)	40 lb.
Casein	10 lb.
Zinc Stearate	5 lb.
Paraffin Emulsion	50-100 lb.
Tarama Emulsion	5 lb.
Formaldehyde (40%)	2 lb.
Triethanolamine	2 lb.
Water	2 10.

The colored design is printed on this paper by using a dye ink having a composition similar to the following:

Acid Dve	100 lb.
Acetone	300 lb.
Divinyl Resin	150 lb.
Methyl Alcohol	100 lb.
Dibutyl Phthalate	10 lb.
Castor Oil	10 lb.

To assist the transfer of the colored pattern to the silk fabric it is advantageous to have present at the time of pressing a volatile solvent which is capable of dissolving the dye but not the coating composition. For assisting the transference of acid dyes to silk fabric it is found that a satisfactory solvent consists of:

318 01:	
Alcohol (95%)	80 gal.
Acetic Acid (36%)	10 gal.
Water	10 gal.

It is claimed that owing to the resiliency of the rubber coating composition and the special manner of applying the transfer paper to the silk fabric, it is possible to obtain very clear and well-graded impressions on crepe materials.

PHOTOGRAPHY

PHOTOG	RAPHY
Fixing Baths Acid Fixing Bath Metric Avoirdupois Water 4 1, 128 oz. Hypo 1160 g. 38 oz. Potassium Metabisulphite 100 g. 3½ oz. The metabisulphite should be added only when the hypo solution is cool, not when it is hot. Chrome Alum Fixing Bath Solution 1 Metric Avoirdupois Water 2½ 1, 80 oz. Hypo 960 g. 2 lb. Sodium Sulphite (Anhydrous) 65 g. 2½ oz. Water to make 3 l. 96 oz. Solution 2 Metric Avoirdupois Water (About 150° F.) 1 l. 32 oz. Potassium Chrome Alum 60 g. 2 oz. Sulphuric Acid C.P. 9 cc. ½ oz. Add solution 2 slowly with constant stirring to solution 1. Acid Hardening Fixing Bath Solution 1 Metric Avoirdupois	A fresh bath should be prepared frequently, as the gelatin-coated backs of the films are likely to become stained in an old or discolored fixing solution. The following Replenisher for two-liter solution of above fixing bath is recommended in cases where the acidity needs to be renewed: Metric Avoirdupois Water Sodium Sulphite (Anhydrous) 15 g. ½ oz. Acetic Acid (28% Pure) 48 cc. 1½ oz. Special Fixing Bath for Printon and Reprolith Films Accuracy in registration for multicolor work being of prime importance, for use in such cases a fixing bath without hardener, as follows is recommended: Metric Avoirdupois Water 1 1. 32 oz. Hypo 485 g. 16 oz. Potassium Metabisulphite 75 g. 2½ oz. In case this bath should lose its acidity by frequent use, giving the film a yellowish stain, add more potassium metabisulphite to restore the acidity of the solution.
Water 4 l. 128 oz. Hypo 960 g. 2 lb.	Acid Hardening Fixing Bath U. S. Patent 1,981,391
Solution 2	Formula No. 1
Metric Avoirdupois	Sodium Thiosulphate 300 g.
Dissolve chemicals thoroughly in order given. Cool solution 2 after mixing and	Acetic Acid 15 cc.
add it slowly with constant stirring to solution 1.	Potassium Alum 15 g. Glycol Borate 10 g. Water to 1 i.

				OMIT III	- 2	:58
	No. 3			Sodium Sulphite		
Sodium Thiosul	phate	300	σ.	(Desiccated) 3 oz.	90 g	
Sodium Sulphit		,	ь.	Water to make 32 oz.	1	
(Desiccated)		15	g.			•
Boron Triacetat		15	g.	Fine Grain Develope	r	
Potassium Alun Water	n	15 to 1	g.	Formula No. 1		
	No. 4	10 1	١.	Avoirdupo	is Mati	rio
Sodium Thiosul		300		Elon 29 gr.	2 g	
Sodium Sulphit		15	K•	Sodium Sulphite	- 8	;•
Acetic Acid	-	15		(Desicented) 3 oz.	100 g	۲.
Citric Acid		1 (g.	145 gr.		
Potassium Alum	1	15 g	ζ.	Hydroquinone 73 gr.	5 g	
Boric Acid Water		to 1	ζ.	Bornx (Crystals) 29 gr. Water to make 32 oz.	2 g 1 f.	
	No. 5	10 1 1	•	No. 2		
Sodium Thiosul		300 g		•	40	
Sodium Sulphite		5 g		Sodium Sulphite p Phenylenediamine	60 g.	
Acetic Acid	•	20 8	c.	Acetone	10 g. 10 ec	
Sodium Acetate		20 g	ζ.	Sodium Metasilicate	3 g.	
Potassium Alum	1	20 g	ζ.	Metol	2 g.	
Borax		20 g	ζ.	Glycin	2 g. ke 2 l.	
Water	No. 6	to 1 l.	•			
		200 ~	_	This is developed for 15 m 65° to 70° F.	inutes	at
Sodium Thiosulp Sodium Sulphite		300 g		65° to 70° F.		
hydrous)	(2111-	15 g		*		
Sodium Acetate	(An-	6	,	Pyrocatechol Developer wi	thout	
hydrous)	•	20 g		Sulphite		
Boric Acid		5 g	.	Pyrocatechol 3- Water 10	4 g. 0 cc.	
Sulphuric Acid (Con-	5 c	.		0 drops	
centrated) Alum				For contrasty negatives use		
Water		15 g to 1 l.		a (Above)	10 cc.	
					100 cc.	
Elon-Hydrogo	uinana Des	cloner		Sodium Carbonate So-		
	Solution	ciopor	- 1	lution (3-4%)	5 cc.	•
Stock	Avoirdupo	a Motr	ic	The second second second		
T21	-			Developer for Film and F	aper.	
Elon Sodium Sulphite	45 gr.		,	Adurol	2 gr.	
(Desiccated)	1½ oz.	45 g	.	Sodium Sulphite	8 gr.	
Hydroquinone	175 gr.		;.	Sodium Carbonate	8 gr.	,
Sodium Carbona	te	0 7 F	.	Water	l oz.	
(Desiccated)	21/4 oz.	67.5 g		Add not more than 1/2 grain	iched	ım An-
Potassium Bro- mide	27 gr.	1.9 g	.	bromide to each ounce of fin veloper. With developer at 70°	F. fil	ms
Water to mal		1 1.	.	will develop in 4 minutes. T	uma G	28
Dilute 1 part to		ter for t	ise.	paper should be exposed so		
Dilute 1 part to			1	image will appear in 45 secon	ds. T	'he
p-Phenylencdi	anine Dev	eloper	1	print will be fully develope	м 111	2
p Phenylene-						
diamine	145 gr.	10 g	.	The times for other papers as		•
Sodium Sulphite		50 g	. 1	Velour black-image appears ute; developed in 2½ minutes.	in i m	ın-
(Desiccated)	1 oz. 290 gr.	50 g	1			1/
Water to mak		1 l.	- 1	Bromide papers—image appea minutes; developed in 3 minut	us III I	74
			- 1	Warmer tones can be obtained		đi-
p-Phenylenediami	ne-Glycin I	Develope	r	luting the developer and givin		
p Phenylene-			- 1	exposure.		
diamine	145 gr.	10 g 12 g		This developer will not affect subject to aniline poisoning. It	perso	TLS
Glycin	175 gr.	12 g	. 1	envices so amine poisomity. Il	, JAMI	i (Till)

quite rapidly and should be kept in a tall, narrow vessel between prints in order to reduce the amount in contact with the air to a minimum.

Gold toner:

Stock Solution

15 gr. 1. Gold Chloride Water

2. 5% Solution Thiourea (1 oz. to 20 oz. water)

For use take 4 drams of gold solution, 3 drams thiourea solution, 5 or 6 drops sulphuric acid and one quart of water. Proceed as follows:

Dilute the required amounts of both stock solutions with one pint of water. Pour gold solution into thiourea solution slowly with stirring. Add the acid to the combined solutions.

Compensating Developer with Pyrogallol

Formula No. 1		
Water	100	cc.
Pyrogallol	0.3	g.
Potassium Metabisulphite		-
(10%)	3	cc.
Caustic Soda (10%)	2	cc.
No. 2		
Water	100	cc.
Pyrogallol	0.3	g.
Potassium Mctabisulphite		•
(10%)	12	cc.
Caustic Soda (10%)	5	cc.

Formula No. 1 at 18° C. (5 to 6 minutes) gives a yellow-brown negative. Formula No. 2 at 18° C. (10 to 12 min-

utes) gives a neutral gray negative and developer can be used a second time.

Modified Hub No. 1 Formula for Glycerin Developer

1000 cc. or (Water 1 qt.) Sodium Sulphite 75 g. 25 g. (375 oz.) Glycin Trisodium Phosphate (Mono-

125 g. hydrate) (4% oz.) Potassium

(45 gr.) Bromide 3 g. This stock solution keeps well, even in partially filled bottles. For use with chloride and chloro-bromide papers it is

diluted with 3 parts of water, and with 4 parts of water for bromide papers. With bromide papers it has been successfully used at temperatures up to 90° F. Because of its high alkalinity, prints developed in this formula should be left in the acid-stop bath for at least 15 or 20 seconds before being placed in

the fixer, and the acid-stop bath should be frequently renewed.

Farmer's Reducer

In case of overexposure or overdevelopment, this well-known reducer can be used effectively for clearing. It is easily compounded by making first a 1:4 solution of plain hypo-for example, 8 oz. of hypo dissolved in 32 oz. of waterand adding to this just enough potassium ferricyanide to turn the solution to a lemon-yellow color. Most workers pre-pare the ferricyanide as a 10% solution in advance, for use as needed; others shake a little of the powder directly into The lemonthe plain hypo solution. The lemon-yellow color disappears with use of the reducer, but may be restored by adding more ferricyanide. The stronger the color, the stronger the reducing action, and vice versa. If the reducer is used too strong its action is not so easy to control.

The film may be immersed in the reducer solution, after being soaked in water to assure even action, or, in cases where only local reduction is desired, the reducer may be applied to the moist film with a tuft of cotton, with rinsing during inspection and afterwards.

Reversing Reversible Film

(1) First Developer

Metric Avoirdupois 1000 cc. 32 oz. Water Metol 2 g. 30 gr. Sodium Sulphite (Anhydrous) 30 g. 1 oz. 12 g. 180 gr. Hydroquinone 8 g. 120 gr. Potassium Bromide 18 g. Sodium Hydroxide 1/2 oz. Potassium Sulphocyanate 5 g.

Develop 4 to 6 minutes at 65° F., depending on exposure.

(2) Wash 5 minutes in running water.

(3) Reversing Bath

Water 1000 cc. Potassium Bichromate 5 g. Sulphuric Acid (Concentrated) Normal bleaching time 3 to 6 minutes.

Keep in bleaching bath until negative image is completely dissolved. (4) Wash 5 minutes in running water.

(5) Clearing Bath

Water 1000 cc. Sodium Sulphite (Dry) 50 g. Clear for 5 minutes.

(6)	\mathbf{Wash}	5	minutes	in	running	water.
-----	-----------------	---	---------	----	---------	--------

(7) Expose to Mazda light or diffused daylight.

(8) Second Developer.

Water	1000	cc.
Metol	5	g.
Hydroquinone	6	ġ.
Sodium Sulphite (Dry)	40	ğ.
Potassium Carbonate	40	ğ.
Potassium Bromide	6	g.
Develop 5 minutes at 65°	F.	
0) (1) 4	4	

(9) Short rinse in running water.

(10) Fixing Bath

Water 1000 cc. 300 g. Нуро Potassium Metabisulphite 50 g. Fix for 2 minutes.

(11) Wash for 30 minutes in running water.

(12) Glycerine Bath

1000 cc. Water 20 cc. Glycerin (C.P.) Leave in glycerin bath for 5 minutes. (13) Remove water with a soft chamois. and dry in a current of warm dry

Note: Operations 7 to 13 take place in

white light. Superpan Reversible film can be desensitized before development by immersion in a 1/5000 solution of Pinacryptol Green desensitizer.

Formula "D16" for Chemically Reversing 16 mm. Film

Water (Distilled) 10 gal. 180 gr. Elon Sodium Sulphite 3 lb. -6 oz. 8 oz. Hydroquinone 9 oz. Sodium Carbonate 1 lb. 1 oz. 63 gr. Potassium Bromide Citric Acid 400 gr. Potassium Metabisulphite 2 oz. Develop 7-15 minutes at 65° F.

Intensifying Formulas

On some occasions and for certain types of work it may be found desirable to intensify film negatives. In such instances the following formulas will give best results, being desirable for their freedom from stain as well as their effective intensifying action.

Mercury Intensifier:

Metric Avoirdupois 1 l. 32 oz. Water 150 gr. Mercuric Chloride 10 g. 75 gr. Potassium Bromide 5 g.

Chromium Intensifier:

This formula gives slightly more vigor ous intensification than the Mercury Intensifier above. Prolonged intensification with it, however, leaves the film with a slight ýellow color.

Metric Avoirdupois Water 1 1. 32 oz.

Potassium Ri-

Hydrochloric Acid 6 cc. 1. 1.6 dr.

Immerse negatives in this solution until bleached, wash for 5 minutes in running water, and redevelop in a Metol Hydroquinone developer. The negatives should then be given a 15-minute wash before drying.

Some intensifying solutions have been known to cause a slight blue coloration of the base of the film. While this is not harmful and does not prolong the printing time unduly, if preferred, such coloration may be easily removed as outlined in the formula for Washing and Drying.

Monckhoven's Intensifier:

Solution A Metric dupois Water 11. 32 oz. Potassium Bromide 23 g. % 02. % 02. Mercuric Chloride 23 g. Solution B Avoir: Metric dupois.

32 or. Water 11. 28 g. % OE. Potassium Cyanide 23 g. Silver Nitrate

The silver and the cyanide are dissolved in separate lots of water, and the former added to the latter until a permanent precipitate is produced. The mixture is allowed to stand 15 minutes, and after filtering, forms Solution B.
Place the negative in A until bleached

through; then rinse and place in Solu-tion B. If intensification is carried too far, the negative may be reduced with a weak solution of hypo.

Because of the deadly poisonous char-

acter of this intensifier, it should be used with care and bottles containing it should be suitably marked.

Agfa Mercuric Iodide Intensifier:

Metric Avoirdupois 200 to 20 to 300 cc. 30 Water Mercuric Chloride 100 cc. 10 (2%)Potassium Iodide 25 cc. (10%)2.5 oz. Hypo (10%) 40 cc. 4 02. Part of the mercury solution is added

and then part of the iodide to the wi solution. atinuing until all the mercury and iodide is added to the water.

When solution is clear, add the hypo. Use full strength.

ercury Intensifier

satisfactory two-solution in-This is alterated two solution are also as a state of the content olution A:

Metric Avoirdapois 1 l. 32 oz. Mercuric Chloride 40 g. olution B: Metric Avoirdupois

Potassium Iodide 200 g. 244

Add B to A until the edution clears.
legative the immersed until changed to brown tolor, then washed and reevelous the inetol-hydroquinone deeloper such as Agfa No. 64. The inensiled negatives after a few and but beld by the control of the hould be type a 16-minute wash before

Intensifier, Photographic

Action (2006)
Priting of Control (2006)
Priting of Control (5%) 1 fl. oz. 1 fl. oz.

Sodium Acetate Solu-

tion (7%)

Internal ery Weak Negatives 400 cc. Water Mercuric Chloride 2 g. 6 g.

1 fl. oz.

Potassium Iodide Each of the dry ingredients is dis-solved in one-half of the water and the two solutions are then mixed. A red precipitate will form at first but will again dissolve, a clear solution resulting.
While the negative attains consider-

able and rapid intensification, it becomes badly colored and will not last very long. To avoid this, the negative is placed in a solution of sodium sulphite for a period of 1/2 to 2 hours. It is then washed thoroughly in water.

If the intensification should be too great it may be reduced in a solution of

sodium cyanide.

Toning Formulas

Sepia Tones by Redevelopment:

Sepia tones may be obtained in any print by subsequent treatment after the print is ordinarily finished. The print should be thoroughly washed before treatment to produce a sepia tone. It is then immersed in the bleaching bath (Solution No. 1) for about 1 minute, or until the middle tones of the print are just perceptible. It is next rinsed thoroughly in cold water and transferred to the redeveloper. When original detail has returned and the print is of desired strength (this will take about half a minute), remove print, rinse thoroughly, and harden by immersion for 5 minutes in the Hardening Solution specified for use in connection with the Fixing Bath (the Hardening Solution only-no Hypo). Finally, remove the print and wash for 30 minutes in running water.

No. 1 Stock Solution

(Bleacher)

The No. 1 Stock Solution, which is the political may be made up for either normal pia tones, warm senia tones, or cold sepia tones, as follows:

For Normal Semia Tones:

Metric Avoirdupois ... Potassium Ferris. cyanide (10% 500 cc. 16 Solution) Potassium Bro-

mide (10% Solution) 100 cc. 31/2 oz. 400 cc. 14 Water oz.

For Warm Sepia Tones:

Metric Avoirdupois Potassium Ferricyanide (10%

600 cc. 191/2 oz. Solution) Potassium Bromide (10% 40 cc. Solution)

Water 360 cc. 12 For approximately a 10% solution, take 100 grains to 2 fluid ounces of water or 10 grams to 100 cc. of water.

For Cold Sepia Tones:

Metric Avoirdupois Potassium Ferricyanide (10% Solution 300 cc. 10 OZ. Potassium Bromide (10% Solution) 500 cc. 16 OZ. 1/2 oz. Ammonia (.910) 10 cc. 190 cc. 61/2 OZ. Water

50 cc.

550

No. 2-Stock 8	Bolutio n
(Re-Develor	per)

Metric Avoirdupois

500 cc. 16 oz. Sodium Sulphide 42.5 g. 11/2 oz. Bleaching Bath for Use.

Avoir-Metric dupois

500 cc. 16 oz. Water No. 1 Stock Solution 500 cc. 16 oz.

(Bleacher) Re-Developing Bath for Use.

Avoir-Metric dupois 32 oz. 1 1.

Water No. 2 Stock Solution 118 cc. (Re-Developer)

Important: Be sure to use sodium sulphide, not sodium sulphite, in compounding the re-developer. Also, use clean trays, free from exposed iron spots, espe-Otherwise blue spots may form on prints.

1. Blue To r Tron Bath First dissover.
Potassium Fereicyanide 375 g.

Potamium Bichromate 1/2 g. Water 40 l.

and pour this solution into a second one consisting of: 425 g. Iron Ammonia Alum 500 g. 40 l.

Oxalic Acid

Water The two solutions must be separately ltered and then mixed at ordinary temrature and with vigorous stirring. bey then form a clear yellowish solu-ion without any sign of turbidity, pro-ided the chemicals have been mixed in he correct quantities and with due re-pard to cleanliness. The time of toning varies according to the tone required.

treating the toned films in a sub-nt weak fixing bath, tones of re-markable clearness are obtained. But it must be expressly noted that, in the case of blue-toned films, the fixing bath must not be used until after a most thorough washing, otherwise a reducing action takes place and detail in the picture is eaten out. The films must be well washed after the second fixing.

2. Uranium Toner (Yellow-Brown) Dissolve:

500 g. Potassium Ferricyanide 10 L Water

Potassium Bichromate (1% Solution)

Then add the whole to:

Uranium Nitrate Oxalic Acid

Water

As in the making up of th the two solutions must filtered and mixed at q ture while stirring was used the should be a selewish solution to be revividence to be revivified from tiles to time during long use by addition of oxalic acid. As much as 1000 g. oxalic acid may be added in all, and so 500 g. of the said is dissolved in water, and the solution added in small doses the time to time, this means, staining of the whites, the otherwise takes that after a time, is readily

avoided. 3. Copper Toning (Reddish-Brown)

Copper Sulphate Sodium Citrate

OF Potassium Citrate Water

To the above add:

Potassium Ferricyanide Water

Potassium Bichromate (1% Solution)

In making the this both separate solution carefully and well mixed at the temperature.

The following observations apply to the use of Baths Nos. 1 to 3.

As is well known, a solution of junction with hypo, ac viz.: the Farmer reducer. Thus a nim which contains only traces of hypo, on being introduced into the toning bath, undergoes a reducing process along with the toning which is aimed at. There are also two conditions which should be invariably observed if it is desired to carry out the toning processes successfully and to keep the toning baths in good conditions:

1. For all toning processes—and this applies also to tinting—frames should be kept for these operations only; frames which have been employed for the development or fixation of prints should on no account be used.

2. Positive film which is to be toned must be especially well washed. In order to ensure that this is the case and to be certain the film is in the necessary state of uniformity, it is advisable to wash the film for a further few minutes immediately before toning.

Wet Collodion Continuous Tone Negative
Plain Collodion 10 g.
To the above add 1 g. of following:
Alcohol 1 l.

Cadmium Iodide
Ammonium Iodide
Cadmium Bromide
Calcium Chloride (6120)

Calcium Chloride (6120)

Respectively.

Re-development increases the opacity if done before fixing and increases the contrast if done after fixing.

Prevention of Haze in Prints German Patent 594,712

The formation of haze is prevented and a blue-black tone imparted to the prints, by adding triazole or tetrazole solution to the emulsion layer or to the developer. Thus, 0.5 to 5 cc. of a 1:100 benzotriazole solution is added to a usual metol-hydroquinone developer.

Control of Photographic Contrasts

M.	
Potash Metabisulphite	160 gr.
Metol	160 gr.
Soda Sulphite	3/4 oz.
Potash Bromide	25 gr.
Water	to 10 oz.
, Q.	
Potash Metabisulphite	160 gr.
Hydroquinone	160 gr.
Soda Sulphite	% oz.
Potash Bromide	40 gr.
Water,	to 10 oz.
* A.	
Soda Carbonate	6 oz.
Water	to 20 oz.

These are concentrated solutions that will keep indefinitely if properly compounded and are diluted for use. In the M and Q solutions the potash metabisulphite should be added to about three-fourths of the water first and partially dissolved, it is not necessary that it should be fully dissolved at this stage, just a good shake up to drive off the oxygen from the water, then the metol or hydroquinone added and fully dissolved before the sods sulphite is added. For use the M and O solutions are used.

For use the M and Q solutions are used either separately or in any proportion desired and an equal volume of the A solution added and then diluted with 3 times the volume of water.

For example, for a normal developer take 1 part of M, 4 parts of Q and 5 parts of A diluted with 15 parts of water. The quantity of water can be varied to suit the particular brand of plate in use, some plates will stand twice this quantity of water. It is a matter of experience.

For positives from very flat negatives the Q solution plus A may be used alone, or a small quantity of M such as 1 of M to 10 or 12 of Q. From very hard negatives the M plus A alone can be used or with a small proportion of Q and, of course, the necessary dilution in each case.

With a high proportion of M to Q the image will appear quickly, but will require time to gain sufficient density. With a high proportion of Q to M the image will appear slowly, but gain density more mpidly in proportion so that the total developing time does not vary so much as would appear at first sight.

sight. To those who have to handle this class of work, either for color half-tone or for photogravure, this system of working is recommended and when once mastered it becomes a very adaptable servant.

Re-Etching Half-Tones with Enamel Off

As in all etching, cleanliness and freedom from grease in the plate to be treated is the first consideration, but any cannel still remaining on the dots is to be left. (This applies to the places to be rolled as well as those where the enamel is good.)

A viscid solution of gum and process white is next prepared:

Gum Arabic
Water
Soz
and when required, to every three parts
of this solution, mix one part process

white.

The plate after being rinsed with water to replace the air between the dots is allowed to drain (not dry) and the gum solution painted over the whole burface. The edge of a wooden rule is next wiped or scraped over the surface in such a way that only the thinnest layer of gum is left on top of the dots

leaving the thick gum remaining between them. A word of warning—should the gum become somewhat thin owing to its application to a wet plate the process must be repeated. Also do not put the gum on a dry plate as it would then be impossible for it to replace the air between the dots. After applying the gum it is dried, using as little heat as possible.

A piece of charcoal having on one of its sides a perfectly flat area of about 1 inch, is now required for rubbing the gum off the surface of the plate, and must be used dry. This flat side is put in contact with the gummed surface and with an even and gentle pressure the gum is rubbed away from the whole surface, or if only to be treated locally, from those parts which are to receive the new ink top. It will be found that very little rubbing is required to remove the gum in the high-lights, while this increases somewhat with the strength of the tone. Rubbing is continued until the metal appears bright and clean, removing any enamel that remains on the areas to be rolled at the same time. If this is carried out properly any increase in tone values owing to the rubbing of the charcoal is negligible, and cannot be seen on a graded strip although etched down beside enamel receiving identical treatment. The gum is now remaining at the sides and between the dots untouched, and the powdered charcoal must be lightly dusted off the surface with cotton wool.

It will be noticed that the white gum between the dots is discolored by the charcoal but this does not matter as in other respects it is quite unaffected. At this stage the roller and ink must come under consideration and these contrary to the usual rule are quite easy to prepare and use. The roller used is a good quality composition roller, and the ink is stone to stone re-transfer ink, both ink and roller are the same as used by line metal printers. Thin the ink with a little pure turpentine in the center of the slab and then evenly distribute the ink over roller and slab. The amount of ink when ready for rolling should be such that it is still possible to see the color of the slab through the ink. The condition of the ink should be just tacky. In rolling up the plates no extra pressure is required, the weight of the roller itself usually being found sufficient. When the whole surface of the plate has received an even layer of ink it is dusted over with fine bitumen powder. This dusting must be done lightly and thoroughly with the aid of cotton wool.

The plate should now be soaked in water for about 2 minutes to soften the gum, but soaking only will not bring it away from between the dots, as a certain amount of force is necessary in the form of a spray of water. The spraying can be done by turning the tap on full and putting the thumb in a position so as to make the water into a narrow beam of as much force as possible, and this is directed all over the surface of the plate, dwelling particularly on those parts (if any) where the gum appears somewhat reluctant to leave, such as a strong cross-line tint. Should any difficulty be experienced in cleaning away ink-covered gum from between the dots, the fault can usually be traced to the gism solution being too thin, or to its imperfect application, but in any case do not attempt other means of removing the gum, such as rubbing with cotton wool, as this will certainly weaken the new top.

After spraying the plate is drained and dried off over the gas with gentle heat, making sure that all moisture is removed before burning-in hard. The temperature reached during the fusing or burning-in of the bitumen and ink should be almost sufficient to burn-in enamel. The required temperature can be judged quite easily in copper by the discolora-tion of the metal: it turning from an orange to a bluish color when approximately the temperature is reached. In zinc there is no discoloration of the metal, but one way to assist the judgment is to paint the back of the plate with shellac and when during the burning-in this turns a dark brown shade, the ink is burnt in.

Burning-in operations completed, the plate, either copper or zinc, is ready for etching as soon as it becomes cold, and it can be chalked with magnesia, staged and treated as though the dots had the original enamel top. One precaution is necessary and that is to take care they are not immersed for any length of time in the acetic and salt bath other than that required to remove the magnesia, as this has a weakening effect on the ink. It is better to dispense with acetic and use a weak solution of nitric acid such as 1 part acid to 20 parts water.

When etching is completed it is sometimes found difficult to remove the ink top even though turpentine and a brush is used, in which case a light rubbing with charcoal will be found the most satisfactory.

Photolithographic Deep-Etched Plates

A fine-grained zinc plate is washed with 5% acetic acid and water, then coated with 1000 cc. of water, 133 cc. of photo-engraver's glue, 100 cc. of 20% ammonium bichromate solution, 20 cc. of ammonium hydroxide at 22° Bé. At 30% relative humidity, the exposure is twice as long as at 60%. The sensitized plate keeps 6 hours at 45 to 50% humidity or 24 hours at 40%. After development in

cold water, the plate is treated for 10 to 15 seconds in hydrochloric acid diluted with 200 parts of water, washed, and dried. Before drying, the image may be dyed in a 2% solution of direct black 2N extra concentrated, or oxydiazol black NJEE. Etching the plate in denatured absolute alcohol to which are added 50 cc. of concentrated hydrochloric acid per liter, for 2 minutes, produces a depth of about 0.0075 mm. The plate is rinsed with alcohol, dried, washed out with asphaltum and liquid reversing ink, and talcked. It is then swabbed with water and in 1000 cc. of water, 400 cc. of 10% barium chloride solution, and 50 cc. of 10% sodium hydroxide solution. Removal of the glue image takes from 5 to 10 minutes. This batch is patented in the United States: 60 cc. of 12 to 14° Bé. Gum arabic solution may be added. After washing with water the plate is bathed for 10 to 15 seconds in very dilute hydrochloric acid, then rinsed in hot water. The plate is next gum-etched and sent to the press.

Photoengraving Enamel U. S. Patent 2,000,453

Glue 20 oz., ammonia solution 2 oz., chromic acid 1.5 oz., and alcohol about 64 oz. are used together.

Planographic and Offset Plates British Patent 421,217

Aluminum plates are made anodes in 0.3-5% nitric for 10 to 30 minutes at a current density of 1 to 2 amperes per square decimeter; zinc plates are made the anode in a saturated potassium carbonate solution for 10 to 30 minutes at a current density of 2 to 3 amperes per square decimeter.

Photographic Masking Paste

Glycerin	1 gal. 3 lb.
Whiting	3 lb.
Neutral Soft Soap	1 lb.

Masking paste must be so formulated as to have sufficient solids or bodying agents that it will not flow down or cause breaks in the film; also it must be capable of being brushed on to form a clean sharp edge. The proportion of glycerin must be sufficient to keep the film from drying up under exposure for at least 48 hours.

Photograph Paste

Gelatin (Photo)

4 oz. 16 oz.

Water 16 oz.
Sonk, dissolve on a water-bath, and add when somewhat cooled:

Glycerin Wood Alcohol Mix 1 oz.

Mounting Translite Prints on Glass

Dissolve 1 oz. of gelatin in 6 oz. of boiled water. After the gelatin has been thoroughly dissolved, add 1 oz. chloral hydratc. Apply the solution to the glass with a brush, coating the glass evenly. Then apply Translite print, wet, face side to the glass. Squeegee with a print-roller until all the surplus gelatin has been removed and air-bubbles are all out. Then allow to dry. This formula will withstand heat more than any other starch or glue formulæ.

Photographic Dry-Mounting Tissue U. S. Patent 2,017,144

A paper mounting tissue is coated on both sides with a composition containing low-viscosity nitrocellulose 100, tritolyl phosphate 110-150 and a resin such as shellac 10-200 parts.

Blue for Drawings

Saturate 10 g. of oxalic acid in a little water with ferric hydroxide, filter off excess of ferric hydroxide, add concentrated solutions of 27 g. sodium oxalate and 11.6 g. sodium ferrocyanide, apply the mixture to paper with a brush and dry in a dark room. Develop the prints with dilute hydrochloric acid or sulphuric acid.

Waterproof Coating for Wooden Photographic Trays

Formula No. 1

Methyl Alcohol 500 cc.
Orange Shellae 100 g.
Rosin 25 g.
Venice Turpentine 25 g.
The ingredients are heated on a waterbath until completely dissolved.

No. 2

One part of gutta percha and one part of paraffin are melted together. When cool, this mixture is dissolved in sufficient benzine to make a mixture of paint-like consistency.

Cleaning Porcelain Photo	graphic Trays
Water	100 cc.
Potassium Cyanide	10 g.
Iodine	3 g.
This is a very satisfactor removing stubborn stains.	ory solution for

Flashlight Powder Formula No. 1

Potassium Chlorate 20 g. Powdered Magnesium 10 g.

The potassium chlorate must first be finely pulverized (to avoid spattering on ignition). It is then carefully mixed with the magnesium. It is preferable to mix this in small quantities on a glass plate, as this mixture is very explosive and a pestle and mortar may prove extremely dangerous.

NO. 2	
Powdered Magnesium	10 g.
Potassium Dichromate	10 g.
This powder is designed to	burn fro

14 to % second.

No. 3

Powdered Magnesium 0.8 g. Ammonium Nitrate

The above should be mixed just before using, the ammonium nitrate being kept in an absolutely dry state. This is a very brilliant and ashless powder and the quantity designated is sufficient for good illumination of a room 15 ft. sq.

Magnesium Flashlight Powder German Patent 592,898

Potassium permanganate, potassium nitrate and sulphur are among the ingredients of a new type of magnesium flashlight powder composition which can be ignited without detonation in cartridges through the medium of a percussion cap. 700 to 900 parts of magnesium are admixed with sulphur (10 to 18), potassium permanganate (100 to 140), potassium itrate (70 to 85), magnesia (100 to 160) and wood charcoal (10 to 20) to 30),

PLATING

Plating on Aluminum

The following formulæ for plating nickel on roughened aluminum are recommended by the Aluminum Co. of America:

Grease is first removed from the surface by immersion in a solution containing:

Sodium Carbonate 1 to 3 oz./gal. Trisodium Phosphate 1 to 3 oz./gal. Temperature about 200° F.

The article to be plated is next rinsed in water and then preferably immersed in 5% hydrofluoric acid solution for about 15 seconds to remove the last traces of alkali and prepare for the etching solution.

The etching solution depends on the chemical composition of the metal.

Formula No. 1

For etching commercially pure aluminum use:

Nickel Chloride	36	oz.
Hydrochloric Acid (sp. gr. 1.18)		gal.
Water Temperature 90° F.	1	gal.

The dipping time should be determined by actual trial. It approximates a half-minute.

No. 2

For etching aluminum alloys containing copper, manganese, and perhaps magnesium use:

Hydrochloric Acid	
(sp. gr. 1.18)	1/3 gal.
Water	% gal.
Manganous Sulphate	1/2 OZ.
Tomporatura 00° F	

The dipping time should be determined by actual trial. It approximates a halfminute.

No. 3

For etching aluminum castings use:
Nitric Acid (sp. gr. 1.42) 3 fl. oz.
Hydrofluoric Acid
(48-52%) 1 fl. oz.

Temperature 75-80° F.

The dipping time should be determined by actual trial. It approximates a halfminute. The container for this etching solution should be lead lined and coated with the following mixture:

Beeswax 1 oz. Paraffin 4 oz.

After etching the articles, they should be well rinsed in water, after which they may be plated in a nickel bath of formula given in Volume II.

Anodic Treatment of Aluminum

The aluminum or aluminum alloy is made the anode in a chromic or sulphuric acid solution, and 10-100 amperes per square foot is passed through for 10-20 mnutes.

Formula No. 1

The chromic acid solution contains 5-15% chromic acid. The current density for this bath varies from 10 amperes per square foot. The temperature of this bath is important and should be kept between 90-100° F.

Fumes of chromic acid develop as the process continues. A ventilating system should be in operation at all times as the fumes are injurious.

No. 2

The sulphuric acid method consists of anodizing the aluminum or its alloy in a solution containing 5-60% sulphuric acid by volume. The current density varies from 10 to 25 amperes per square foot. The temperature control is not as important as in the chromic acid solution.

Sulphuric acid spray is released during the process, and for this reason the bath should have a ventilating system applied to it.

After the work has been removed from the solution, it is essential to wash with water until all traces of sulphuric acid or chromic acid have been removed. For this purpose two rinses in running water for 10 minutes each will suffice.

> Anodic Coating of Aluminum Formula No. 1

British Patent 427,308

The electrolyte consists of an acid to which a glucoside or hydrolyzed glu-

coside has been added. A suitable bath consists of 100 l. sulphuric acid of sp. gr. 1.220 to which is added 300 g. baptisin or 500 g. hydrolyzed barbaloin. Alternatively, 500 g. trihydroxymethylanthraquinone as obtained from the hydrolysis of frangulin may be added.

No. 2 British Patent 429,344

a . . . a . 1-

Caustic Bous		zu g.
Water		20 g. 1 l.
Glycerin		150 cc.
In place of the following may be	glycerin any e used:	one of the
Formaldehyde		75 cc.
·	or	
Lactose		90 g.
	or	

Barbaloin 50 g. Operate at 10-15 volts; current density 18-24 amperes per square foot at 15-25° C.

Coloring Aluminum

If anodized aluminum is placed in a solution of an organic dye, the dye unites with the coating formed on the aluminum and forms a colored lake. These colors will not wash out. Thus, by dipping anodized aluminum in a green dye solution, a green coating is obtained. In this way any desired color can be obtained.

Formation of Noncorrosive Film on Aluminum, Magnesium or Their Alloys

Japanese Patent 109,261

Aluminum, magnesium or their alloys are boiled in a solution of 25 g. of ammonium molybdate and 25 g. of ammonium tartrate per liter.

Antimony Plating

Antimony Oxide	60 g.
Hydrofluoric Acid	114 g.
Water	1000 cc.
Aloin	1/4 g.
Clovel Oil	1/8 g⋅

The mixture should be stirred until solution of the oxide is complete. A lead vessel can be used. Vessels of these materials or of wax can be used as containers for the final plating bath. Wax vessels cannot be used in the making of the bath due to heat of the reaction. A cast antimony anode is used. This bath must be electrolyzed for several days.

perhaps to eliminate impurities, before good deposits can be obtained.

A current of 0.8 ampere per sq. dm. (7.4 amperes per sq. ft.) can be used. Higher currents give less smooth deposits. Deposits can be made any thickness even 1 cm. (0.4 in.) or more. The current efficiency at the cathode is practically 100%.

Brass Plating

Copper Cyanide	4.2	oz.	per	gal.	
Zinc Cyanide	1.5	OZ.	per	gal.	
Sodium Cyanide	6.7	oz.	per	gal.	
Sodium Carbonate	4	0 Z.	per	gal.	
Ammonium Hy-			•	•	
Sodium Carbonate	4				

droxide 0.12 oz. per gal.
Use brass anodes and 2-4 amperes per square foot.

Bronze Electroplating Bath British Patent 412,277

Copper Cyanide	40 g.
Sodium Stannate	20 g.
Sodium Cyanide	35 g.
Caustic Soda	5 g.
Water	to make 1 l.

Brass and Bronze Solutions

Brass Solution:

Copper Cyanide	4 oz.
Zine Cyanide	1 oz.
Sodium Cyanide	6 oz.
Sodium Carbonate	2 oz.
Water	1 gal.

Temperature 90° F. Cathode current density 2.5 to 3 amperes per sq. ft.; 2 to 3 volts. Use rolled anodes, 80% copper, 20% zinc.

Bronze Solution:

Copper Cynnide	4 oz.
Zinc Cyanide	1/2 oz.
Sodium Cyanide	5 oz.
Sodium Carbonate	2 oz.
Rochelle Salts	2 oz.
Water	1 gal.

Temperature 95° F. Cathode current density, 2 to 2.5 amperes per sq. ft.; 2 to 3 volts. Rolled bronze anodes, 90% copper, 10% zinc.

Cadmium Solution:

Sodium Cyanide	9 oz.
Cadmium Oxide	3 oz.
Caustic Soda	2 oz.
Water	1 gal.

Temperature 80° F. Cathode current density, 8 to 10 amperes per sq. ft.; 2 to 2½ volts. Use iron and cadmium anodes,

one iron to three cadmium. cadmium anodes when solution is not in

Cadmium Plating Bath Formula No. 1

Cadmium Oxide 3 oz. per gal. 10 oz. per gal. Sodium Cyanide

No. 2

Cadmium Oxide	39.4 g.
Potassium Cyanide	128.2 g.
Sodium Sulphate	50 g.
Nickel Sulphate	1 g.

Cadmium-Zinc Alloy Plating

Satisfactory deposition is possible from solutions containing 55-75 g. of zinc, 5-30 g. of cadmium, 3-6 mg. of gelatin or caffeine, and 15-20 g. of aluminum sulphate per liter, operated at 25° with pH 4 and current density 1-2 amperes per square decimeter. The cadmium content of the alloy is increased by rotating the cathode and raising the temperature and is decreased by raising the current density, increasing the acidity, and using addition agents and salts. Complex organic nitrogen addition compounds, e.g., caffeine and aloin, have a selective effect, retarding cadmium deposition and thus permitting the cad-mium of the bath to be increased. Alloys containing 45-55% of zinc show most resistance to corrosion by aqueous sodium chloride.

Cadmium Plating Die Castings

Scratch brush raw die casting wet or if rough, polish first. Articles are then cadmium plated and given either a dry or wet scratch brush for desired finish. Lacquer to protect finish. Satisfactory deposits may be obtained from the following solution:

Sodium Cyanide 7 oz./gal. Cadmium Oxide 3 oz./gal. Potassium Hydroxide 2 oz./gal. Temperature 113° F.

Current density 10-25 amp, per sq. ft.

Any patented brightener may be used. Strip, 10% ammonium nitrate.

Chromium Plating

The chromic acid salt to be used should consist (according to British Standard Specification) of

Chromium Trioxide (CrO₃) 99.5 % Sulphate (as Sulphuric Acid) 0.2 %

0.05% Chlorides (as Chlorine) Insoluble Matter 0.15%

and the solution made up of 250-500 g. per liter, with a density of 25 to 27° Bé. Sulphate is added in a proportion of 1/100th of the chromic acid concentration; with too high amount of sulphate, current and throwing power fall off badly. Fluoride may be substituted for sulphate, calcium fluoride 30 g./l. in a 500 g./l. solution gives good results. The solutions should be made up very carefully; usually the bath works best when aged artificially. The tank for the solution (of glass, wood, lead-lined metal) should be arranged for heating as temperature is a critical condition; 40° C. (100° F.) is usually applied, sometimes 60° C. (140° F.) may be required, while for thick, dull deposits cold solution can be used.

Anodes are of lead or lead-antimony alloy; the latter is less affected when the bath is not operating. Current density is very important; for bright deposits on nickel 150 amperes per sq. ft., for thick deposits 300-400 amperes per sq. ft. are used. The high current density requires a particularly careful suspension of the work in the bath, thin wires as in other plating practice are out of the question; very often special jigs are used. In cer-tain cases, where the work is rather large, auxiliary anodes of lead or iron are arranged to insure a good deposit inside a hole, recess, etc. Degreasing in trichloroethylene, polishing and nickel-plating before chromium plating is desirable. Careful subsequent treatment is essential to avoid corrosive effects of eventually remaining bath solution; repeated rinsing alternately in hot and cold water, drying in an oven or hot sawdust is necessary.

Chromium Plating Bath

Chromium Oxide (Free from Sulphuric Acid) 350 g. Potassium Fluoride 3 g. Water 1000 cc. Run at 18-20° C., using 3.8 to 4 volts.

Chromium Solutions Formula No. 1

Chromic Acid 33 oz. Sulphuric Acid 0.3 oz. Water 1 gal.

Total sulphate (from both the chromic acid and the sulphuric acid) should be

Temperature 113° F. Cathode current density 125 to 1750 amperes per sq. ft.

No. 2		
Chromic Acid	5 5	oz.
Sulphuric Acid	0.55	0 Z.
Water	1	gal

Total sulphate (from both the chromic acid and the sulphuric acid) should be 0.55 oz.

Temperature 95° F. Cathode current density 75 to 125 amperes per sq. ft.

The anodes and temperature control coils should be of 6% antimonial lead. The chromic acid tanks should be of steel, lined with 6% antimonial lead.

No. 1 is used where heavy deposits are desired.

No. 2 is used where the deposit is for decorative purposes.

Cobalt Plating Bath British Patent 427,458

Cobalt Chloride		40-150	
Sodium Acid Fluoride		10-40	
Ammonium Chloride		15-60	
Cobalt Basic Acetate		15-60	g.
Water	to	make 1	ſ.

Copper Solutions

No.	1
31/2	
41/2	oz.
2	oz.
1/14	0Z.
1	gal.
	No. 31/4 41/4 2 1

No. 2		
Copper Carbonate	5	oz.
Sodium Cyanide	10	oz.
Hyposulphite of Soda	1/64	
Water	1	ga

Either solution should be operated at 100° F. to 110° F. Cathode current density 4 to 6 amperes per sq. ft.; 1½ to 2 volts. Use rolled copper anodes.

Acid Copper Solution

Copper Sulphate	28 oz.
Copper Sulphate Sulphuric Acid Water	3 to 5 fl. oz. 1 gal.

Temperature 75° F. Cathode current density for still solution 10 to 15 amperes per sq. ft.; % to 1 volt. Agitation of the cathode or of the solution allows the use of higher current density. Use rolled copper anodes.

Coppering by Immersion

Copper Sulphate Sulphuric Acid Water		to	1	oz. oz. gal.
--	--	----	---	--------------------

Where only a very thin film of copper is desired the above solution will give good results.

Acid Copper Plating

Cupric Sulphate 27 oz. per gal. Sulphuric Acid 7 oz. per gal.

Use brass anodes and a current density of 20-40 amperes per sq. ft.

Cyanide Copper Plating

Copper Cyanide
Sodium Cyanide
Sodium Carbonate
Sodium Carbonate
Sodium Carbonate
Soz. per gal.
Use at 35° C. at 13 amperes per sq. ft.

Blue Dip (for Plating Copper and Brass Articles)

Bichloride of Mercury	⅓ oz.
Sodium Cyanide	6 oz.
Ammonium Chloride Water	1 oz. 1 gal.
TTALCI	T Rai.

Fluoride Bath

A 41	g.		
Antimony Oxide (Commercial)	60	8	OZ.
Hydrofluoric Acid			
(Commercial,			
48%)	114	15.3	OZ.
Water	1000	1.6	gal.
Aloin	0.25	0.033	oz.
Clove Oil	0.012	0.0016	OZ.

The last two constituents, the so-called addition agents, are used up during the plating; hence, they must be added regularly to the bath. The quantities given above are sufficient for about 12 hours of operation.

Immersion Gold Solution

Fulminate of Gold	4	dwt.
Yellow Prussiate Potash	24	oz.
Carbonate of Soda	12	0 Z.
Caustic Soda		0 z.
Water	1	gal.

Solution should be boiled in a cast iron tank for an hour and allowed to cool to 180° F. before using.

Salt Water Gold

Dair marci Gold	
Yellow Prussiate of Potash	64 oz.
Sodium Phosphate	32 oz.
Sodium Carbonate	16 oz.
Sodium Sulphite	8 oz.
Gold as Fulminate	12 dwt.
Water	4 gal.
Solution is boiled for one	hour, then

diluted with water to make 4 gal. of solution. The solution is placed in a porous pot which is put in a tank that contains a saturated solution of sodium chloride heated to 190° F.

Green Gold

Metallic Gold as Fulmi-		
nate or Cyanide		4 dwt.
nate or Cyanide Silver Cyanide		1/4 dwt.
Sodium Cyanide		2 oz.
Carbonate of Soda		2 oz.
Water		1 gal.
Temperature 105° F.;	2	volts; 18
karat green gold anodes.		

Rose Gold

Yellow Prussiate of Potash 4 oz.
Potassium Carbonate 4 oz.
Sodium Cyanide 14 oz.
Gold as Fulminate 10 dwt.
Water 1 gal.
Temperature 175° F.; 6 volts. If a red color is desired, add small quantity of copper carbonate.

Coating Iron with Lead and Tin

Iron and steel can be coated electrolytically after pickling with sulphuric acid, in a bath of tin borofluoride, lead borofluoride and borofluoric acid with acid-proof layers of a lead-tin alloy which are so elastic that the metals can still be worked mechanically; the temperature must, however, not rise above 150 to 200° C, as otherwise the coatings would melt. The deposits are made at a current density of 0.5-3.0 amperes per sq. dm.

Electrolytic Burnishing of Iron

Oxidize anodically in 20 to 40% caustic soda at 1 to 6 amperes per sq. dm. at 1 to 2 volts at 60-70° C.

Thin Deposits of Iron

Dissolve 16 oz. of ammonium chloride in each gallon of water. Connect up tank, same as for plating, using cold rolled iron for anodes. On the cathode rod suspend some old plating racks or other work, and work solution with highest current density obtainable. After 4 or 5 hours of work of the solution, there will be enough iron dissolved from the anodes and the solution will produce a deposit of irom. Operate solution at 80° F.; 1.5 to 3 ampages per sq. ft.; 1 volt.

Iron Solution

Ferrous Chloride 40 oz. Calcium Chloride 20 oz. Water 1 gal.

Temperature 200° F.; current density 40 to 50 amperes per sq. ft.; 2 to 2½ volts; pH 1.5 to 2; pure iron anodes.

volts; pH 1.5 to 2; pure iron anodes.

This bath is used to produce heavy deposits of iron.

Preparing High-Speed Steels for Plating

In order to secure good adhesion of electro-deposits to high-speed steel it is treated anodically at 2.7 amperes per sq. dm. in a bath containing 115 g. of caustic soda and 15 g. of citric acid per liter until gas evolution is uniform over the whole surface, then rinsed with water, dipped momentarily in 6-12N-hydrochloric acid and finally washed with water.

Electrodeposition of Lead

Fifty grams of lead perchlorate and 10 g. perchloric acid in 1 liter electrolyte and a current density of 0.25-0.50 amperes per square decimeter are recommended for the preparation of pure 0.1 mm. deposits of lead of good texture. Agitation of the bath permits a higher current density and thicker deposits. Addition of 0.2-0.4 g. peptone and moderate agitation improve the deposit and allow a current density of 1 ampere per square decimeter. Higher current densities up to 2 amperes per square decimeter require constant and efficient stirring and heating up to 60° C. permits 3-4 amperes per square decimeter. For technical purposes 1 ampere per square decimeter is recommended.

Lead Solutions

Lead Carbonate	20	oz.
Hydrofluoric Acid (5	0%) 32	oz.
Boric Acid	14	oz.
Glue	0.025	oz.

Place the hydrofluoric acid in a leadlined tank and add the boric acid with constant stirring. When the boric acid is completely dissolved, the solution is allowed to stand until cool, when the lead carbonate is added in the form of a paste with water. The solution is allowed to settle in the plating tank. The solution is then diluted to the proper volume with water and the glue added after dissolving the same in warm water. Mechanical agitation of the solution is ressential.

A cathode current density of 10 to 20 amperes per sq. ft., 3 to 4 volts, and lead anodes are employed.

Thin Deposits of Lead

Carbonate of	Lead	2 oz	
Caustic Soda		6 oz	
Water		1 g	ıl.
Lead anodes.	Temperature	175°	F.

3 to 4 volts.

Coating Magnesium and Its Alloys French Patent 766,685

Magnesium or an alloy thereof is coated by introducing it into a rotating drum along with an alloy of zinc (25) and cadmium (75 parts) and some gal-vanized iron turnings. The drum is heated to about 290° C., when the alloy becomes pasty, and is rotated for about 3

Commercial Nickel Plating

The three principal methods of nickel plating, i.e., ordinary plating in the sta-tionary bath, rapid plating and barrel plating are discussed and compared as to their respective economic advantages. In all methods it is necessary that new nickel sulphate be continuously formed at the anode and that the deposit be fine in grain. The deposit must permit of mechanical working without injury. The deposit if chromium plated must not peel. The composition of an ordinary stationary bath consists of 75 g. nickel ammonium sulphate in one liter water with a pH of about 5.8; increasing the latter to 6.4 increases, reducing it to 4.6 decreases the throwing power of the bath. Specific gravity is 6-7° Bé., the current density 0.3 ampere per square decimeter, voltage 3.5, temperature 18° C. A thickness of 0.025 mm. is obtained in 7 hours. A rapid plating bath must work at 50°, the grain of such deposit is the finer, the better the electrical conductivity of the bath. The compositions used are: 240 g. nickel sulphate, 30 g. boric are: 240 g. nickel sulphate, 30 g. noric acid, 19 g. potassium chloride in 1 liter water; or 240 g. nickel sulphate, 120 g. magnesium sulphate, 30 g. boric acid; or 240 g. nickel sulphate, 30 g. boric acid; or 240 g. nickel sulphate, 30 g. boric acid, 150 g. magnesium sulphate, 10 g. sodium chloride, 50 g. sodium sulphate, 0.1 g. sodium fluoride in 1 liter water. The current density must be adapted to the kind of ware to be plated. Pure nickel anodes do not dissolve as easily as 98% nickel anodes. If the deposition velocity is too high, an excess of oxygen is formed at the anode, passivates it and finally nickel bisulphate and peroxide are formed without nickel going into solu-tion. Plating in the barrel requires a pH of not less than 6.6, at 8-12 volts, time usually 2 hours, bath temperature 35-50°.

Nickel Solutions

Nickel Solution for Brass, Copper, and Cold Rolled Steel

A nickel solution that has been used with good results on brass, copper and cold rolled steels is made as follows:

Formula No. 1

Double Nickel Salts	8 oz.
Single Nickel Salts	4 oz.
Boric Acid	2 oz.
Sodium Chloride	2 oz.
Water	1 gal.

Solution to be operated at 80° F.; 2 to 21/2 volts; 6 to 8 amperes per sq. ft. and a pH of 5.8.

For solutions that are operated at a higher temperature and a correspondingly higher current density, use:

No 2

Double Nickel Salts	8 oz.
Single Nickel Salts	8 oz.
Sodium Chloride	3 oz.
Boric Acid	3 oz.
Water	1 gal.

Temperature 110° F.; 21/2 to 3 volts; 20 amperes per sq. ft., and a pH of 6; depolarized nickel anodes 99% plus. Replenish by the addition of single nickel an lta.

Low pH Solution for Heavy Deposits of Nickel

No. 3

Single Nickel Salts 32 c Sodium Chlorido 6 c	
T) 1 4 17 A 4	Z.
Boric Acid 4 0	Z.
Water 1	zal.

Nickel Strip

Bulphuric Acid		4 OZ.
Water		1 oz.
Temperature 800	F . lood	eathodes.

volts. If 3 or 4 oz. of copper sulphate per gallon are dissolved in the water before adding to the acid, the strip will not attack the base metal so readily.

Nickel Brighteners

Bright deposits of nickel are obtained from No. 1 formula above by the use of cadmium chloride or one of the prepared brighteners that are on the market. The pitting of nickel deposits is siminated by adding hydrogen peroxide to the 12th Use from 1 to 5 cc. of 100 within peroxide to each gallon deposits appear the severity of the pitting

Nickel Plating

The nickel content of the bath is about 40-50 g. per liter; current density 0.3-0.4 amperes per square decimeter while for rapid plating methods 1-3 amperes per square decimeter are employed. The bath is stirred and the pieces are moved to avoid streaks on the deposit, pH is 5.8-6.2. For rapid nickel plating the following bath is recommended: pure nickel sulphate 22.5 kg., pure ammonium sulphate 2.0 kg., pure nickel chloride 0.5 kg., pure sodium perborate 0.5 kg., water 100 liters 35-40° C., voltage 2.75-3.5.

Hydrogen Poor Nickel Plating

Nickel sulphate 80 g., nickel fluoride 8 g., sodium chloride 1 g., sodium sulphate 0.5 g., sodium nitrate 0.02 g., sulphosodium-phenolate 0.12 g., sodium citrate 2 g., boric acid 6 g., zircon-ammonium fluoride 0.2 g., all in 1 liter water. The ammonium fluoride binds the hydrogen and the deposits adhere well to the base. The voltage employed with this bath is 2 volts.

White Nickel Plating Formula No. 1 (Low Metal Bath)

Nickel Sulphate
Ammonium Chloride
Boric Acid
pH = 5.4

12 oz. per gal.
2 oz. per gal.

Use at room temperature with nickel anodes, and 10-20 amperes per sq. ft.

No. 2

(High Metal Bath)

Nickel Sulphate
Nickel Chloride
Boric Acid
pH = 5.3

134 oz. per gal.
4 oz. per gal.

Use nickel anodes and a current density of 15-45 amperes per sq. ft. with a temperature of 50-60° C.

Nickel Bath for Die Castings

Nickel Sulphate
Ammonium Chloride
Boric Acid
Sodium Sulphate
Bodium Citrate
pH = 5.5

pH = 5.5
Temperature, 20-30° ; current denaity = 15-30 amperes perseq. ft.

Depositing Nickel on Rough Steel

If a smooth deposit is required over
rough steel, instead of buffing down the

steel, it is possible to pickle the steel in an acid until all the scale is removed and then depositing a heavy coat of copper, using an acid sulphate bath for this purpose. The heavy coat of copper is then buffed until it is smooth. The coat can now be finished in any way desirable. It is much cheaper to buff copper than steel,

Black Nickel Plating

Nickel Ammonium
Sulphate
Sodium Sulphote
Sodium Sulphocyanate
pH = 5.8-6.0

60 g. per l.
14 g. per l.

Gray Nickel Plating

Nickel Ammonium
Sulphate
Sodium Sulphocyanate
pH = 5.4

60 g. per l.
14 g. per l.

Plating Zinc with Nickel

(1) Strike for 5-10 minutes in any suitable cold nickel solution. The following formula is suggested:

Nickel Sulphate 15 oz. per gal. Anhydrous Sodium

Sulphate 15-18 oz. per gal. Ammonium Chloride 2-3 oz. per gal. Boric Acid 2 oz. per gal. Temperature 78-85° F. pH = 4.9-5.4 (electrometric)* Current density 24-30 amp. per sq. ft.

(2) Rinse thoroughly in cold water.

(3) Transfer without drying to the following solution:

Nickel Sulphate 20 oz. per gal. Ammonium Chloride 4 oz. per gal. Boric Acid 2 oz. per gal. Temperature 105-115° F.

pH = 5.0-5.3 (electrometric) Current density 40-80 amp. per sq. ft.

* May be increased to as high as 30 ounces per gallon for intricate shapes.

Solvent Cleaning of Zinc

Grease and oil may be removed from zinc and zinc alloy castings by the use of trichloroethylene, carbon tetrachloride, xylol, ethyl acetate, etc. These solvents are most effective when used in apparatus involving vapor rinsing. However, these solvents do not remove oxide films and zinc salts and hence where parts are to be electroplated, the metal should subsequently be submitted to an

acid dip which serves the additional purpose of roughening the surface to provide good adhesion of the finish coating. The following solutions have been used in zine alloy die castings:

(1) Phosphoric acid etch—treat for 30 seconds in 3% solution of phosphoric acid (85% H₃PO₄ grade, specific gravity 1.74) rinse and dry.

(2) Hydrochloric acid etch—treat for 30 seconds in a 10% solution of hydrochloric acid (35 to 37% HCl grade, specific gravity 1.18-1.19) rinse and dry.

(3) Hydrofluoric acid etch—treat for 30 seconds in a 1% solution of hydrofluoric acid solution (48% HF grade) rinse and dry.

Plating of Zinc

Considering nickel and nickel-chromium plated coatings on zine and zine alloy eastings, a minimum thickness of coating of 0.0003 in. at the thinnest point is necessary to give any satisfaction in outdoor service. Completely satisfactory quality will not be obtained consistently with coatings of less than 0.001 in. average thickness.

Nickel Plating Solutions Formula No. 1

Nickel Sulphate 10 oz. per gal. Anhydrous Sodium

Sulphate 10-15 oz. per gal.
Anmonium Chloride 2-3 oz. per gal.
Boric Acid 2 oz. per gal.

Operating details for this solution follow:

pH—This should be held between 5.3 and 5.7 electrometric or 5.8-6.2 colorimetric. The anode area should be controlled to minimize pH changes. pH should be checked daily and adjustments made by the addition of ammonium hydroxide or sulphuric acid as needed. Under best operating conditions this solution will tend slowly to become alkaline.

Temperature—For use in applying nickel directly on zinc this solution should be kept at or preferably slightly above room temperature (70-80° F.). If the temperature falls below 70° F. the deposits will be hard and brittle showing cracks. Temperature above 80° F. will tend to cause the formation of black streaks in recesses.

Nickel Content—The prescribed nickel sulphate content corresponds to about 2 oz. per gallon of nickel calculated as metal. No harm will result if this increases somewhat in use. Sodium Sulphate Content—The amount of sodium sulphate present in the solution should be regulated to suit the complexity of the articles to be plated. Simple shapes may require not more than 10 oz. per gallon of sodium sulphate. More complicated shapes may require the presence of 15 oz. per gallon or more. Some commercial platers add as high as 30 oz. per gallon. In general, the sodium sulphate content should be the lowest possible for the articles being plated.

Current Density—When made up according to the formula given, the bath should be operated at hetween 12 and 20 amperes per sq. ft. The maximum current density will be determined by the tendency for the deposits to burn. In the presence of very high sodium sulphate concentrations, burning may develop at current densities lower than 20 amperes per sq. ft. If streaking occurs at the maximum current density, purification of the solution may be necessary.

Agitation—Agitation reduces porosity and permits the use of somewhat higher current densities. With certain shapes, agitation will be found absolutely necessary for successful plating.

Pitting—Like all other nickel solutions this bath will at times develop a tendency towards pitting. This is usually an indication that foreign matter is present. A temporary cure can be effected by adding hydrogen peroxide or sodium perborate to the solution. Permanent freedom from pitting can only be obtained by continuous filtration and scrupulous care in avoiding the presence of foreign material in the solution. Pitting nay on occasion develop from faulty cleaning.

No. 2

Nickel Sulphate
Anhydrous Sodium
Sulphate
15 oz. per gal.

Sulphate 15 oz. per gal.
Ammonium Chloride 3 oz. per gal.
Boric Acid 2 oz. per gal.

Operating details for this solution are given below:

pH—Should be kept between 4.9 and 5.4 - Celetrometric or 5.4-5.9 colorimetric by means of additions of sodium hydride or hydrochlorio acid. Ammonium droxide and sulphuric acid should not be used as the solution is nearly saturated with respect to nickel ammonium sulphate.

Temperature—The more concentrated solution permits the use of somewhat higher current densities which in turn permit the use of higher temperatures of

operation which may be reflected in slightly softer deposits. The minimum safe temperature is 75° F. and the maximum is 87° F.

Nickel Content—Corresponds to about 3 oz. per gallon calculated as nickel metal. Any large increase in nickel content may result in crystallization of double nickel salts from solution.

Sodium Sulphate Content—Should be regulated as for the 2-oz. (nickel content) solution. In general, somewhat higher sodium sulphate contents will be required in the present case.

Current Density—This more concentrated solution permits the use of higher current densities, the range in the present case lying between 24 and 36 amperes per sq. ft.

Agitation Pitting—The considerations mentioned under Formula No. 1 above hold in the present case.

No :

Nickel Sulphate 20 oz. per gal. Ammonium Chloride 4 oz. per gal. Boric Acid 2 oz. per gal.

Operating details for this solution are given below:

pH—The pH of this solution should be held between 5.0 and 5.3 electrometric or 5.5–5.8 colorimetric. Higher pH will cause cracking and peeling while lower pH will tend to increase the attack of the solution on exposed portions of the base.

Temperature—Should be between 105 and 115° P. (40-45° C.). Lower temperatures will not permit the deposition of soft nickel. Higher temperatures, while allowable, tend to cause excessive loss of water by evaporation.

Current Density—The current density should under no circumstances fall below 40 amperes per sq. ft. and preferably should be maintained at 60 amperes per sq. ft. or higher. Not only does the speed of production fall off at the lower current densities but contamination of the solution becomes more serious. These current densities are similar to those required for chromium plating and suitable guerator capacity should be available.

Agitation—Agitation will tend to reduce pitting and porosity.

Pating—Like most warm solutions, new baths of this composition may develop an exaggerated type of pitting. This condition can be readily overcome by additions of hydrogen peroxide. Sodium perborate should never be used for the reasons given below.

Sodium Salts—Sodium salts should not be permitted to enter! this solution. When the solution is pure, very high current densities can be employed without burning. The presence of sodium salts very definitely restricts the operation to low current densities which not only do not utilize the full production capacity of the solution but also permit excessive zinc pickup. For these reasons the rinsing between nickel tanks should be thorough, sodium perborate should not be used to prevent pitting, and additions of alkali to raise pH should be made with ammonium hydroxide.

Nickel Plating Methods

Three methods of applying adequate nickel coatings to zinc and zinc alloy castings have been found successful.

Multiple Nickel

This method consists essentially of depositing on the zinc articles, from either Formula No. 1 or No. 2 above, a coating of nickel 0.0001 in. to 0.0002 in. thick following which the articles are thoroughly rinsed in cold water and placed in a warm nickel solution (Formula No. 3) for completion of the plating to the required thickness.

The strike coating must be adequate to protect the zinc base from the action of the subsequently used warm solution. For simple shapes a 5-minute deposit at 25 amperes per sq. ft. may be sufficient. More complicated shapes will need 10 minutes at this current density.

Rinsing—In the interval between the two nickel tanks the articles should not be allowed to dry. If drying does occur poor adhesion of the second coat will develop. The use of cold water in the rinse will minimize the danger of this happening.

Copper-Nickel

While the system of plating nickel direct has a great many advantages, good results have also been obtained commercially by plating with copper-nickel deposits totalling 0.001 in, in thickness.

In this system of plating, the work is cleaned thoroughly, a coating of copper is applied to a thickness of 0.0005 in. from a copper-cyanide solution, followed, after rinsing, by the application of 0.0005 in. of nickel in a warm nickel solution (Formula No. 3).

The copper cyanide solution may be any one of those commonly used. A typical formula follows:

Sodium, Cyanide	4-6 oz./gal.	(30-45	Ø. 1	oer 1.	١
Copper "Cyanide	4 oz./gal.			er l.	
Sodium Bicarbonate					
Sodium Bisulphate	1/4 oz./gal.	(1.87			

The solution should be used at 70-113° F. (21-45° C.) with a current density of 10-15 amperes per sq. ft.

8

The copper-nickel system of plating is adapted to the production of heavy deposits. Its use is not advocated for coatings less than 0.0005 in. in thickness. The copper layer should be at least 0.0002 in. thick in order to avoid complete absorption by the zinc base and to provide protection of the zinc base from attack by the warm nickel solution. The copper layer fills the same role here as the primary or strike nickel deposit in the multiple nickel system of plating.

The nickel deposit must be at least 0.0003 in. thick for outdoor use. Thinner deposits will readily permit the seepage through pores of copper salts which will stain the surface with an unsightly brown

Nickel-Copper-Nickel

When coatings ranging from 0.00075 in. upward are desired, multiple contings are necessary to avoid cracking. Multiple nickel coatings have been described above. The system nickel-copper-nickel has also been used successfully.

Clean the articles thoroughly as described under "Cleaning of Zinc and

Zinc Alloys.'

Plate 0.0002 in. of nickel in either cold solution described in Formulas No. 1 and No. 2.

Plate 0.0004 in. of copper from an acid-copper solution.

Color copper, coat, and clean. Plate 0.0004 in. of nickel from any warm nickel solution such as described in Formula No. 3 above.

The buffing operation is not essential if the two primary coats are sufficiently smooth to make coloring of the final nickel readily accomplished.

The acid copper solution may be of any accepted composition. The following formula is typical:

Copper Sulphate 24 oz./gal. Sulphuric Acid 6-8 oz./gal.

This solution is used at room temperature to 113° F. (45° C.) with a current density of 10-50 amperes per sq. ft. Animal glue may be used as a brightener in amounts of 1/2 oz. per gal. (0.9 g. per l.).

Bright Nickel Plating on Zine

A bright nickel deposit which requires no buffing or coloring can be produced in the sulphate type of solution by the addition of 100 of an oz. per gal. of cad-mium sulphate. A small amount of cadmium sulphate may be added from time to time to maintain the cadmium metal content in use.

The deposits produced are very bright and smooth but somewhat brittle and should not be deformed or bent. Chromium should not be deposited over such coatings as the additional stress will

crack and peel the nickel.

Bright nickel deposits of this type tend to be brittle and are suitable only for use in thin form for indoor application.

Black Nickel Plating on Zinc

A bright, black, adherent coating can be obtained on zinc by a 2-minute plat-ing in the following solution at 113° F. (45° C.).

Nickel Ammonium

Sulphate 8 oz./gal. Zine Sulphate 1 oz./gal. Sodium Sulphocyanate 2 oz./gal. 1-2 amp./sq. ft. Current Density

Chromium Plating * on Zinc

Chromium may be applied either as a thin finish coating over nickel or as a heavy protective coating directly on zinc from the following solutions:

Chromium Oxide (CrO₃) 33 oz./gal. 0.3 oz./gal. Sulphuric Acid (H2SO4)

Chromium Oxide (CrO₃) 33 oz./gal. Chromium Sulphate $(Cr_2(SO_4)_3)$ 0.44 oz./gal.

For finish plating this should be used at 113° F. (45° C.) with lead anodes and at a current density of 75-150 amp. per sq. ft. A 3-6 minute deposit should be sufficient.

For heavy deposits applied directly on zinc these solutions may be used with the conditions of operations stated.

* No consideration has been given to the patent situation involving chromium solutions which must be taken into account by the

work should be plated for 20-25 minutes to insure reasonable thickness of coating. The deposits obtained will not be bright but will have a luster ranging from milky to frosty depending upon conditions. The explanation for the failure to obtain bright deposits apparently lies in the fact that these solutions etch the surface of the zinc slightly before deposition occurs to protect it. The deposits can, if only milky, be readily buffed to a bright Instar.

Somewhat better protection and ease of buffing will be obtained with chromium deposits applied directly on the zinc from these solutions at room temperature with a current density of 50-125 perature with a current density of or local amp, per sq. ft. The deposits will be dull gray in appearance but can be readily buffed or brushed to a high luster. The work should be plated for 20-25 minutes to insure a good protective plate.

Cadmium Plating * on Zinc

Recent practice to improve the surface appearance of zinc alloy die castings such as carburetors, etc., which do not require a fine fluish is to cadmium plate them directly without buffing. Satisfactory deposits may be obtained from any of the numerous types of solution in use. A typical formula is:

of process roundary rou	
Sodium Cyanide	7 oz./gal.
Cadmium Oxide	3 oz./gal.
Caustic Potash	2 oz./gal.

This solution should be used at room temperature to 133° F. (45° C.) with a current density of 10-25 amp. per sq. ft. Almost any of the patented brighteners will give satisfactory results.

* No consideration has been given here to the patent situation involving cadmium plat-ing which must be taken into account by the

Stripping Methods Nickel-Chromium

Chromium and nickel may be removed by making the work anode in concentrated sulphuric acid to which a small quantity of commercial glycerin is added. Zinc is only slowly attacked by the concentrated acid but as the solution absorbs moisture from the air this attack will increase to the point where pitting of the zinc starts and the solution demands attention. The excess moisture may be removed by boiling the solution until heavy white fumes appear.

Nickel Coatings

Immerse in the following cold solu-

Water	1	part
Sulphuric Acid	2	parts
Nitric Acid	2	parts
Hydrochloric Acid	1/16	part

Prepare by adding the sulphuric and nitric acids to water and, after allowing the solution to cool, adding the hydrochloric acid.

Non-Electric Nickel Plating Compound

rormula No. 1	
Nickel Ammonium Phosphate	5 oz.
Nickel Sulphate	3 oz.
Cream of Tartar	2 oz.
Tin Chloride	2 oz.
Ammonium Chloride	1 oz.
Codium Chloride	1 oz.
Copper Powder	2 oz.
Chalk Powder (Whiting or	

4-5 oz. Precipitated Carbonate) until pasty

No. 2	
Nickel Ammonium Sulphate	25 g.
Nickel Sulphate	15 g.
Cream of Tartar	10 g.
Tin Chloride	10 g.
Ammonium Chloride	5 g.
Salt	3 g.
Whiting	20 g.
Metallic Copper, Powder	10 g.
Water until	pasty_

Rhenium Plating

Rhenium, with an atomic weight of 186.3, is a very heavy metal. It is both ductile and malleable, and has a brinell hardness of 250. It is quite soluble in nitric acid but insoluble in hydrochloric acid. Therefore it should find wide use for plating on jewelry, as the hydro-chloric acid released in perspiration will not affect the deposit.

Bath 1

Potassium Perrhenate 11 g. per l. Sulphuric Acid 9.3 g. per l. Temperature, 25°-45° C. (77°-113° F.) Current Density, 90-110 amp. per sq. ft.

Bath 2

Perrhenic Acid	20 g.	per l.
Sulphuric Acid	5 g.	per l.
Temperature, 25°-30° C.	(77°-86°	F.)
Current Density, 90-140	amp. per	sq. ft.

Bath 3

Dissolve 8 g. of rhenium in concentrated nitric acid. Add 4 cc. of concentrated sulphuric acid, and boil until sulphur trioxide fumes are evolved. Dilute to one liter, and add enough sulphuric acid until 6 g. per l. is obtained. This solution may be used at 20°-60° C. with 50-100 amp. per sq. ft. using platinum as an insoluble anode, or rhenium as an anode. The metal deposits as a smooth shiny adhering deposit. The plating time can be 10-60 minutes.

Rhenium Nickel Plating

Potassium Perrhenate
Nickel Sulphate
Sulphuric Acid
Temperature, 25°-50° C.

Current Density, 50-60 amp. per sq. ft.

The alloy of nickel rhenium obtained from the above solution is somewhat

lighter in color than pure rhenium.

Rhodium Plating

Five g. of rhodium chloride in 1 l. water are boiled with 40 g. of sodium nitrite until light yellow; 3 g. of sodium carbonate are added to remove traces of his muth and the solution is filtered. After cooling 50 cc. of saturated aqueous ammonium chloride are added and precipitated ammonium rhodinitrite is collected and washed with cold water. 8.52 g. are heated to fuming with 33 cc. of concentrated sulphuric acid cooled, and diluted to 11. Deposition is best effected at 40° C. with platinum electrodes using a current density of 5 amp. per sq. ft. Cathode current efficiency is about 45%.

Rhodium Plating Silver Canadian Patent 343,808

Five g. of rhodium ammonium nitrate is dissolved in 1 l. of boiling water containing 20 cc. of sulphuric acid, and after the reaction is completed 100 g. sodium nitrate are added. The mixture is evaporated to dryness and the residue dissolved in 1 l. of water to form an electrolyte for plating silver. Deposition is preferably conducted at 80-100° F. with a current of 20-50 amp. per sq. ft. of cathode surface and an inert anode, such as carbon or platinum. The plated silver resists tarnishing.

Non-Poisonous Silver Plating

TION F ORDOROUS	 	
Silver Nitrate	25-30	
Thiourea.	60-70	g.
Water	1	l.

Use 0.2 amp. per sq. dm. at 30-35° C. at 1½ volts-

Silver Dip

Silver Chloride 1¼ oz./gal. Sodium Cyanide 2½ oz./gal.

In order to apply this procedure to headlight reflectors it is necessary to remove any nickel plate, then polish and clean before dipping. The film of silver so produced is very thin and will have a short useful life.

Improving Silver Finish

There is no bright dip for silver in the same way as a dip for brass or copper. The surface of the parts in question can be improved by making them anodes in a solution containing 8 oz./gal. of sodium for cyanide and 8 oz./gal of sodium ferrocyanide. Use 10-15 amp./sq. ft. and about 6 volts pressure. Keep work well agitated.

Non-Poisonous Silver Plating

Citric Acid 60 g.
Sodium Iodide 520 g.
Use a silver anode with current density
of 1-1.8 amp. per sq. dm.

Silver Plating Stainless Steel

In silver plating stainless steel it is essential to etch slightly the surface with an acid pickle. This is done to obtain a metallic surface that the subsequent electro-deposit of silver will adhere to

A pickle made up of 10%-15% sulphuric acid, either electrolytic or still, at a temperature of 150°-160° F., will work satusfactorily.

A silver plating bath of the following composition can be used:

Silver Cyanide 4 troy oz./gal.
Sodium Cyanide 5 oz./gal.
Free Cyanide 4 oz./gal.
Water 1 zal.

Non-Electric Silver Plating Compound
Silver Nitrate 6 oz.
Ammonium Chloride 6 oz.
Sodium Thiosulphate 10 oz.
Calcium Carbonate or Chalk 10 oz.
Water until pasty

Brightener for Silver Solution

Silver Solution Sodium Cyanide	1 qt. 8 oz.
Carbon Bisulphide	1 oz.
Ether	1 oz.

To prepare the brightener place the carbon bisulphide and ether in a quart

bottle and shake thoroughly. Dissolve the cyanide in the silver solution and fill bottle. Shake bottle from time to time until the carbon bisulphide is thoroughly dissolved and then filter. One ounce of this stock solution should be sufficient for an addition to each 50 gal. of the regular plating solution. Care must be taken to avoid an excess.

Silver Strips

	F'ormula.	No.	1		
Sodium	Cyanide			12	oz.
Caustic				2	oz.
Water				1	gal.

Reverse current with cold rolled steel as cathodes. Voltage 6 to 8. Agitate the work for a cleaner job.

No. 2

Nitric Acid	1 gal.
Place crock that contains ot water container. If all	

Place crock that contains the strip in a hot water container. If all water is kept from the strip, brass or copper work will be attacked only slightly.

Removing Fire Scale from Silver
Nitric Acid 2 oz.
Water 1 oz.
Use hot and agitate work.

Removing Fire Scale by Reverse Current Sodium Cyanide 8 oz. Water 1 gal. Use hot and agitate work. Lead anodes; 4-6 volts.

Bright Dip

2011 But - T	
Sulphuric Acid	2 gal.
Nitric Acid	1 gal.
Water	1 qt.

Add 1 oz. of muriatic acid for 5 gal. of above.

It is necessary to add water only when a new bright dip is made. Dip must be operated cold.

Matt Dip

Sulphuric Acid	1 gal.
Nitric Acid	1 gal. 2 lb.
Zinc Oxide	2 lb.

Operate hot and keep out all water and chlorides. If the matt is coarse, add sulphuric; if too fine nitric acid.

Gold Solutions

Cyanide Solution

 Metallie Gold as fulminate or Cyanide
 5 dwt.

 Sodium Cyanide
 2 oz.

 Sodium Phosphate
 1 oz.

 Water
 1 gal.

 Temperature 130-160° F.; 1 volt; 24

Temperature 130-160° F.; 1 voit; 24 kt. gold anodes.

Chloride Solution

Gold Chloride	6	oz.
Hydrochloric Acid	10	oz.
Water	1	gal.
Room Temperature; 2-3	volts.	

In preparing the solution dissolve the gold chloride in dilute hydrochloric acid before adding it to the solution.

Silver Solution

Formula No. 1

Silver Cyanide	31/2	oz.
Sodium Cyanide	5	oz.
Sodium Carbonate	2	oz.
Water	1	gal.
No. 2		
Silver Cyanida	314	07

Silver Cyanide 3½ oz.
Sodium Cyanide 8 oz.
Sodium Carbonate 2 oz.
Water 1 gal.

Either of the two solutions will give good results if operated at a temperature of 75° F. with a cathode current density of 4 or 5 amp. per sq. ft.; ¾ to 1 volt. Formula No. 1 is generally used, but the deposit of No. 2 is whiter.

Silver Strike

Silver Cyanide % oz.
Sodium Cyanide 8 oz.
Water 1 gal.
Use steel or carbon anodes; 6 volts.

Black or Gun Metal Finish on Steel A black or gun metal finish may be obtained on steel articles by heating them in a retort with a small amount of charred bone and heated to 700°-800° F. After articles are thoroughly oxidized temperature is dropped to 650° F. and a mixture of bone and bone oil is added. Several hours are required to produce finish. Articles after coming from retort are rolled in oily granulated cork until uniform black finish is secured.

The following solution will give to aluminum a uniform black color:

Water 1 l. Potassium Permanganate 5-10 g. Nitric Acid 28° Bé. 2- 4 cc. 20-25 g. Copper Nitrate Temperature, 80° C.

Time to obtain deep black, 20-30 minutes.

Tantulum Plating U. S. Patent 1,933,319

The electrolyte is a fused mixture of Potassium Chloride 120 g. Potassium Fluoride Potassium Tantulum Fluoride 100 g. 25 g. Tantulum Oxide

in a graphite crucible at 750° C. This bath gives a bright plate on iron or nickel at 1 to 10 amp. per sq. dm.

Tin-Plating from An Alkaline Bath

Tin-plating of copper, brass, zinc, lead, hard lead, iron, steel and aluminum can best be carried out at 0.15-0.5 volt in alkaline aqueous stannous chloride, or in alkaline aqueous sodium stannate plus sodium chloride, with 0.12-0.2 g. of gelatin per l. A tin anode (anode current density 0.45-1.6 amp. per sq. dm.) can be used. A cathode current density is 0.2-1.5 amp. per sq. din. The maximum and minimum concentrations of the bath are 50 g. of tin salt for 2 molecules of sodium hydroxide and 12 g. for 1 molecule respectively.

Non-Poisonous Tin Bath

An alkaline tin bath without cyanides to be used at 50-60° C. is composed of sodium stannate 7.5 kg., sodium acetate 1.25 kg., sodium hydroxide 1.25 kg., starch 70 g., water 100 l. Anodes are partly of tin, partly of iron. The bath can be used for electrical tinning of kitchen utensils.

Tin Solution

1111 001441011	
Sodium Stannate	12 oz.
Caustic Soda	1 oz.
Sodium Acetate	2 oz.
Hydrogen Peroxide	1/12 oz.
(25 Volume) or	
Sodium Perborate	⅓ oz.
Water	1 gal.
The relation is spended	of a femner

The solution is operated at a temperature of 140-160° F.: 4 to 6 volts; anode current density, 20-60 amp. per sq. ft.

Immersion Tin Solution

Tin Chloride	1/2 oz.
Aluminum Sulphate	2 oz.
Cream Tartar	2 oz.
Water	1 gal.
Water	I gai.

The solution is allowed to boil for 30 to 45 minutes and the addition of a very small quantity of sulphuric acid (about 1 drop to each gal. of solution) hastens the deposition of the tin deposit.

Caustic Soda Method (Tin)

This method is used to tin by immersion, small brass or copper articles.

Caustic Soda	12 oz.
Stannous Chloride	4 oz.
Sodium Chloride	1 oz.
Water	1 gal.

The solution is placed in an iron tank, which is heated with a steam coil. The bottom of the tank is covered with moss tin over which is placed an iron wire screen. The work to be tinned is bright dipped or tumbled clean, placed in brass wire baskets and separated with sheets of perforated tin, placed in solution at boiling temperature for 15 to 30 minutes, or until covered with tin. Rinse thoroughly in clean cold water, hot water, dry in sawdust.

Protecting Tin and Lead Against Corrosion

French Patent 777,314

Dip in following solution:

Copper Sulphate	25 g.
Nickel Sulphate	15 g.
Ammonium Molybdate	3 g. 1 l.
Water	1 I.

Tungsten Plating

The Carbonate Bath:

Tungstic Acid 125 g. per l. 330 g. per l.

Sodium Carbonate 330 g. per L'se at 90° C., 50 amp. per sq. ft.

The Phosphate Bath:

100 g. per l. Tungstic Acid Sodium Phosphate

500 g. per l. (Na₃PO₄·12H₂O)

Use at 90° C, with 50 amp. per sq. ft.

Citric Acid Bath:

100 g. per l. Tungstic Acid Potassium Hydroxide 70 g. per l. Citric Acid 250 cc. per 1.

(2.5 Molar Citric Acid)

Use platinum anodes; 50 amp. per sq. ft. at 20° C.

Electrolytic Surface Treatment of Zinc

British Patent 421,696

Zinc and alloys consisting mainly thereof are provided with an insoluble

coating resistant to weathering and corrosion by anodic treatment in a substantially neutral electrolyte containing an alkali metal ferrocyanide, ferricyanide, dichromate, oxalate or molybdate or ammonium oxalate or molybdate or more than 1 of these. Suitable baths contain 35 g. crystal ammonium oxalate or 50 g. crystal potassium ferrocyanide per l. The metal surface may first be cleaned by cathodic treatment in a bath containing 45 g. sodium phosphate, tribasic, per l. The coatings may be painted, lacquered or dyed, color coatings being obtainable by adding a dye to the electrolyte.

Zinc Solutions Acid Zinc Solution

Zinc Sulphate	32	oz.
Ammonium Chloride	2	oz.
Sodium Acetate	2	oz.
Water	1	gal.

Temperature 80° F. Cathode current density, 15-20 amp. per sq. ft.; 3-4 volts; pH, 3.5-4.5, using thymol blue as a indicator.

Cyanide Zinc Solution

Zinc Cyanide	4 oz.
Sodium Cyanide	4 oz.
Caustic Soda	3 oz.
Water	1 gal.

Temperature 100° F. Cathode current density 10-15 amp. per sq. ft.; 2-3 volts; keep free cyanide equal to metal content. Use pure zinc anodes. Finish work by rinsing in cold water, then hot water, then drying in hardwood sawdust.

Zinc Cadmium Alloy Plating Zinc Sulphate 295 g. per l.

Zinc Sulphate 295 g. per l. Cadmium Sulphate 50 g. per l. Aluminum Sulphate 30 g. per l. Caffeine or Licorice 5 mg. per l.

Sulphuric acid may be used in small amounts, but as a general rule, the deposit will not be as bright if acid is present, although appreciably harder. This alloy coating can be deposited directly upon iron, steel, brass, bronze, copper, etc.

Coloring Zinc Dark Brown U. S. Patent 1,853,323

Zinc or die cast zinc can be colored dark brown by treating in a bath containing:

Chromic Acid 200 g. per l. Sulphuric Acid 2 g. per l. provided the material is treated with an alternating current.

Cleaner for Barrel Plating

Water	1 gal.
Soda Ash	6 oz.
Caustic Soda	2 oz.

This is not suitable for work which has soldered or tinned parts. Such parts should be cleaned in a cleaner which does not readily attack solder or tin. This should be used, 8 oz. to each gal. of water. More may be used without any bad effect upon such work immersed not more than 20 minutes, which will ordinarily clean almost any "hard to clean" parts. It is understood of course that the solution should be kept hot, 180° F.

This cleaner does not readily tarnish brass and copper and has a considerable amount of insoluble material in it which has a scrubbing effect when boiling. This is very effective also in removing oils and dirt and does not require frequent replenishing.

This cleaner is sold on the market under various trade names, the only difference being in the proportions of the 3 sodium compounds.

Another effective cleaning solution used hot or boiling is composed as follows:

Water	l gal.
Soda Ash	4 oz.
Caustic Soda	2 oz.
Trigodium Phognhata	2 0=

This too may be varied to suit almost any requirement in cleaning, but a solution made up weaker than the above formula will not work well long. The formula approximates very closely many proprietary cleaners now on the market.

One of the best and simplest combinations for an electrical cleaner is as follows:

ows:	
Water	1 gal.
Soda Ash	2 oz.
Caustic Soda	1 oz.
Trisodium Phosphate	1 oz.

This may be modified to meet almost any problem of cleaning with the current.

Cleaning Enamel from Metals

Using 50 amp. per sq. ft. at 2½ volts, reversing polarity at 10-second intervals and using following bath gives excellent results.

Caustic Soda

13.6 oz.

Trisodium Phosphate 6.38 oz. Sodium Silicate 1.62 oz. Water to make 1 gal.

Cleaning Phosphor Bronze Sheets

After the regular sulphuric acid pickling, they are treated in a bath made of a 10% solution of sulphuric acid with 14 to 1/2 lb. of sodium bichromate added to each gal. of the solution.

The general practice is to heat the solution with live steam.

The quantities given are for each gal. of water in the cleaning tank. Have the water near the boiling point and add the materials by dusting on the surface and stirring until dissolved.

Electroplating Radiators British Patent 425,846

Copper cynanide 40, sedium stannate 20, total sedium cyanide 65, sedium hydroxide, 7.5 g. per l. is specified, this having a free sedium cyanide centent of 20 g. per l. and pHI 13. A current density of 1-80 amp. per sq. ft., or higher, and a bath temperature 15-17° C. are used. A deposit containing 13-16% tin is obtained. A suitable alloy for automobile radiator shells is tin 15% and copper 85%. The anode preferably consists of an alloy in the proportions of the desired deposit but these may vary by 10% or more. The anodes should be heat-treated to obtain a maximum softness by casting in a metal mold, cooling in the mold,

heating to 1000° F. for 15 minutes and quenching in water. The alkalinity of the lath should be maintained at a pH 12.8-13.5 and the free sodium cyanide at 10-45 g. per l.

Coloring Razor Blades Blue

After blades have been hardened and drawn and being sure that surfaces are absolutely clean, polish well and heat to 550-600° F. This temperature will not affect temper.

Protection of Magnesium by Means of Selenium Coatings

Of many methods tried for conting magnesium with selenium, the following give the same results; (1) immersion for 3 hours in an aqueous solution of 8% sodium selenite, 3.2% selenious acid and 0.10% sodium chloride at 80-90° C; (2) a 10% selenious acid solution with 0.1-0.5% sodium chloride for 5-10 minutes; (3) a 2% sodium selenite solution with 0.2% phosphoric acid for 1 minute; (4) initial cleaning for 30 seconds in 1% chromic acid at 80° and then treatment as in method 3; (5) cleaning as in method 4 followed by method 2.

Increasing Life of Graphite Electrodes

To increase their resistance to attack during electrolysis anodes 25 × 25 mm. in size are soaked in coal tar for 1½-2 hours at 150-180° F., or in pitch for 3-5 hours at 300-350°. They are then heated at 300-500° to drive out the more volatile compounds. Larger anodes require longer treatment. Such anodes are more stable and more efficient than anodes treated with linseed oil. Mixtures of tar and pitch, or bakelite lacquers, may also be used.

POLISHES, ABRASIVES

Aluminum Polish	. c. Calcium Carbonate,
Formula No. 1	Precipitated 20 oz.
Potassium Hydroxide 40 g.	Iron Oxide—Red 5 oz.
Water 900 cc.	d. Ammonia (10%) 4 oz.
Olive Elaine 150 cc.	e. Alcohol 10 oz.
Alcohol 25 cc.	Dissolve a in an enameled or zinc
Ethylene Dichloride 50 cc.	plated or tin-plated steam-heated kettle;
Add the potassium hydroxide to th	
water, warm to 75° C. and slowly stir i	form homogeneous emulsion, then add the powders c. Saponify with d, let
the olive Elaine until completely dissolved. Cool and add the alcohol an	stand several hours, and add e. Then
solved. Cool and add the alcohol an	impregnate rags in this solution.
ethylene dichloride.	-
Directions for Use	Non-Scouring Copper Polish
Dip a piece of fine steel wool or roug	Moke a posts of feely namidened along
cloth into a liquid and rub on to the al	and mineral oil This will not constab
minum. Then wash the surface with he	/C
water and dry as usual. This aluminut	
polish used in dish water in proportion of about 2 tbsp. per 1 gal. will softe	
the water and assist in cleaning.	b. Water, Hot 53 cc.
<u>-</u>	L c. Olein, Distilled 5 cc.
No. 2	
Whiting 75 g.	e. Alconol, Denatured 16 cc.
Tripoli, Fine, Yellow 20 g. Sodium Bicarbonate 3 g.	f. Tripoli 20 g.
Potassium Sulphocyanide 2 g.	Dissolve a and b, saponify with c, di
Add Glycerin Water (25%) unt	lute with e, add f.
pasty.	
	Chromium Polishes
Silver Plating Polish	Formula No. 1
(Renews as it polishes)	Olein 20 cc.
Silver Nitrate 30 oz.	Stearin 60 g.
Salt 30 oz.	Melt.
Cream of Tartar 200 oz.	Calcium Carbonate
Grind and sift through 100 mesh sieve	e. (Powdered) 20–30 g. Cool, powder.
Then make into a paste with	Cool, powder.
	No. 0
"Cellosolve" 50 parts	No. 2
Water 50 parts	Chromium Oxide 60 g.
Water 50 parts	Chromium Oxide 60 g. Stearic Acid or
Water 50 parts Silver Polish	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g.
Water 50 parts Silver Polish Soap 20 oz.	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3
Water 50 parts Silver Polish 20 oz. Stearic Acid 5 oz.	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g.
Water 50 parts Silver Polish Soap 20 oz. Stearic Acid 5 oz. Gilders Whiting 32 oz.	Chromium Oxide 60 g. Stearic Acid or 40 g. Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g.
Water	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Japan Wax 15 g.
Water 50 parts Silver Polish 20 oz. Stearic Acid 5 oz. Gilders Whiting 32 oz. Tripoli 3 oz.	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Japan Wax 15 g. Paraffin Wax (46-48° C.) 60 g.
Silver Polish Soap 20 oz.	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Japan Wax 15 g. Paraffin Wax (46-48° C.) 60 g. Melt on water bath.
Water	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Japan Wax 15 g. Paraffin Wax (46-48° C.) 60 g. Melt on water bath. Melt together and add:
Water	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Paraffin Wax (46-48° C.) 60 g. Melt on water bath. Melt together and add: Turpentine 130 cc.
Water	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Japan Wax 15 g. Paraffin Wax (46-48° C.) 60 g. Melt on water bath. Melt together and add: Turpentine 130 cc. Tripoli, Dry 70 g.
Solver Polish Soap 20 oz.	Chromium Oxide 60 g. Stearic Acid or Paraffin Wax 40 g. No. 3 Carnauba Wax 10 g. Yellow Wax 15 g. Japan Wax 15 g. Paraffin Wax (46-48° C.) 60 g. Melt on water bath. Melt together and add: Turpentine 130 cc. Tripoli, Dry 70 g.

POLISHES, ABRASIVES 285		
No. 4	Black Polish for Over	ns
Rouge (Iron Oxide) 50 g.	Formula No. 1	
Kieselguhr, White, Burned 100 g.		1000 lb.
Neuburger Chaik 150 g.	Graphite, Flaky	50 lb.
Coconut Oil Soap 700 g.	Lampblack	10 lb.
No. 5	Beeswax, Crude Montan Wax, Crude	100 lb.
Chromium Oxide, Powdered 50 g.	Paraffin Scales (50-52° C.)	
Paraffin Wax 50 g.	Melt together.	
Emery 30-50 g.		5 lb.
No. 6	Nigrosine, Fat Soluble	til pasty
Stearin 90 g.		til pasty
Stearin Oil 25-30 cc.	No. 2	
Neuburger Chalk 30-45 g.	Graphite, Colloidal Paraffin Wax	20 lb.
Melt together.	Paraffin Wax	13 lb.
Cool; powder.	Lacquer Benzoline (White	07 IL
	Spirits)	67 lb.
Polish for Metals	No. 3	
	a. Olein, Distillate	15 cc.
French Patent 772,648	Stearin (52-54° C. titer)	5 g.
Formula No. 1	b. Ammonia (25%)	4 cc.
A polishing compound contains kaolin	c. Spindle Oil	10 cc.
30-50, talc 10-20, rosin 18-30, alcohol	Alcohol	40–50 cc.
4-15, ammonia 5-18 and acetone 1-10	Melt a on water bath, say	onify with
parts by weight.	b, add c, then an abrasive (1	mery, Car-
No. 2	borundum, Chromium Green,	Graphite).
French Patent 772,691		
A compound contains powdered silicon		
dioxide 35, soap powder 5.9, neutral oil	Automobile Polish Cle	aner
dioxide 35, soap powder 5.9, neutral oil 0.23, ammonium sulphate 3.1 and ben-	Formula No. 1	
tonite 0.63 kilograms.	f Olein	10 cc.
No. 3	Mineral Oil	20 cc.
Kieselguhr 2 parts	a. Petroleum	20 cc.
Strong Ammonia Water 1 part	Turpentine Oil or	00
Denatured Alcohol 1 part	White Spirit	28 cc.
Shake well with water q.s. to give	Alcohol	6 cc.
creamy consistency.	b. Ammonia (0.910)	6 cc.
	c. Infusorial Earth	10 g.
Metal Polish (Sidol Type)	No. 0	
a. Olein, Distilled 4.5 cc.	No. 2	10 oz.
Stearin 1 g.	Yellow Wax *Air-Floated Tripoli	18 oz.
Alcohol 5 cc.	White Spirit	19 oz.
Heat to 50° C.	Soft Soap	1/2 oz.
b. Ammonia (sp. gr. 0.91) 7 cc.	Water	21/2 oz.
Saponify.	Melt the wax in a double	ha bus use
	the powder slowly; keep st	irring while
c. Oxalic Acid 2 g. Water (50-60° C.) 70 cc.	slowly adding the white spir	it. Dissolve
d. Neuburger Chalk 25 g.	the soft soap in the water	and add to
Optional:—add more water.	the mix with constant stirring	g. On cool-
Optional:—add more water.	ing this forms a soft paste.	
Metal Polish Block	A liquid polish can be n	nade as fol-
	lows:	
a. ?	No. 3	01/ mt
01014	White Spirit	21/2 pt. 21/2 pt.
b. Spindle Oil, Refined 2-10 cc. Vienna Lime 30 g.	Mineral Oil Turkey Red Oil	4 pt.
c. English Red (Ferrous	Ammonia	1 oz.
Oxide) 38-30 g.	Water	5 pt.
Mix first b, to prevent saponification	Glycerin	1 pt.
of the fats a	Formaldehyde	8 oz.
	-	

Fuller's Earth Bentonite	8 6	0Z.
Dontomico		

Mix the oils together first and add the abrasive powders, then the water, ammonia, glycerin, and formaldehyde; stir rapidly until a smooth mixture is obtained.

*The quantity and type of abrasive used can be varied according to whether the polish is to have a strong or mild abrasive action. Pollshes to be used as maintenance polishes by car owners should be only mildly abrasive, otherwise too much of the finish will be rubbed off.

Car Polishes Formula No. 1

a. Spindle Oil, Refined Methyl Hexalin	80-85 20-15	
b. Distilled, Warmed Water	400-900	g.

Add b to the mixture a with high speed stirring.

Apply spraying and polish with a rag.

No. 2	
Linseed Oil	200 g.
Dipentene	300 g.
Paraffin Oil	200 g.
Petroleum, Refined	250 g.
Camphor Oil, Light	50 g.
Apply simply with rag.	

Automobile Cleaner and Polish

Kieselguhr	30 oz.
Tripoli	5 oz.
Paraffin Wax	4.5 oz.
Carnauba Wax	0.5 oz.
Varnolene	30 oz.
Tint with iron oxide.	

Automobile Paste Polish

Carnauba Wax	5 oz.
Beeswax	5 oz.
Ceresin Wax	5 oz.
Stearic Acid	2 oz.
Soap	2 oz.
Varnolene	45 oz.
Water	10 oz.

Automobile Polish, Powdered

Mineral Oil	5 lb.
Kerosene	10 lb.
Diglycol Laurate	1 lb.
Silica Dust	⅓ lb.
Kieselguhr	4 lb.
Tripoli	1 lb.
Tithom	

Automobile Polish (Tumbler's) U. S. Patent 1,969,387

To 3½ gal. of pale blown castor oil, add ¾ gal. of orthodichlorbenzol. This is mixture No. 1. To 15 gal. of water, add 11 gal. of neutral pale mineral oil and ¾ gal. of ammonia, which has been previously made up of one part of ammonia of 26° Bé. and 4 parts of water. This is mixture No. 2. Mixture No. 1 and mixture No. 2 are combined and agitated for about 5 minutes. Three and one-half gallons of special petroleum spirit is added and the whole mass is now stirred about 10 minutes. It is then run through a colloid mill and is ready for use. Alternatively, all of the ingredients may be mixed in a single batch and passed through the colloid mill, which breaks up the particles to a fine degree. This obviates preparing separate mixtures.

Auto Polish U. S. Patent 1,979,787 Wax Base

Carnauba Wax	66.5 g.
Petrolatum Wax (160 to	·
165° F. Melting Point) Petrolatum (140° F. Melt-	26.6 g.
Petrolatum (140° F. Melt-	
ing Point)	6.3 g.
Rosin	0.6 g.
Wax Base (Prepared as	
Above Described)	9 g.
Refined Mineral Oil (Nar-	
row Cut)	41 g.
Starch	41 g. 0.5 g.
Water	49.5 g.
	. 0

The refined oil is a distillate having an initial boiling point of about 350° F. and an end point of about 475° F. Although it is not necessary that these precise limits be maintained, it is important that a narrow cut be used of about this range. The so-called "W.W. 150" (water white kerosene), with a boiling range of about 373 to 504° F. evaporates too slowly, while oleum spirits, with a boiling range of about 300 to 425° F. evaporates too rapidly to give best results. The narrow boiling range of the refined oil is of particular importance in a "set" or solid emulsion of this type. It is also of particular importance that the oil be highly refined (treated with sulphuric acid for the removal of unsaturateds and other impurities) because untreated light petroleum distillates may be injurious to the skin.

In preparing the finished product melt

the base stock with the refined oil and heat the mixture to a temperature of about 175 to 200° F. Then boil a 1% starch solution and make an oil-in-water emulsion in a colloid mill at a temperature above the melting point of the wax and below the boiling point of the water, usually at about 130 to 200° F. When the resulting emulsion cools, it sets to form a semi-hard, solidified emulsion which is extremely stable and which possesses entirely different structural properties from the ordinary liquid oil inwater emulsions of the same concentrations. The product may be stored for an indefinite period of time without separation, and it may be easily handled and applied.

Solid Abrasive Polish (Wax), Automobile Formula No. 1 a. Montan Wax, Bleached Paraffin (40-42° C.) 8 g. 8 g. 2 g. Ozokerite, Refined 35 g. Infusorial Earth b. Spindle Oil, Refined White Spirit 13 cc. 13 cc. Turpentine Oil or 21 сс. Substitute No. 2 Montan Wax, Bleached 8 g. Montan Wax, Double 5 g. Bleached 2 cc. Olein Potassium Carbonate 2 g. 3 cc. b. Glycerin (28° Bé.) 40 cc. Water, Boiling c. Yellow Clay or to suit Bentonite Turpentine Oil or 22 сс. White Spirit Melt a, add hot (boiling) b, then c; cool, add d.

Auto Polish

Formula No. 1

	g.
Paraffin Wax (50-52° C.) 5	g.
Hard Soan	g.
	cc.
Water Soluble Dyestuff 2	g.
(Black: 4 parts Nigrosine)	
(Black: 4 parts Nigrosine) Ammonium Hydroxide (0.910) 1	cc.
Alcohol 20	cc.
No. 2	

Montan Wax, Bleached	7	g
Soft Soap	3	
Potassium Carbonate	0.8	8

Water Water Soluble Dyestuff (Black: 4 parts Nigrosine)	87.2 2	cc. g.
No. 3		
Shellac (Orange) Alcohol Carnauba Wax Paraffin Wax (50-52° C.) Turpentine	2	g. g. g.

Polish for Lacquered or Polished Objects Quine Detent 170 724

owiss ratent 1/2,	130	
Turpentine	100	cc.
Paraffin	50	g.
Beeswax	15	g.
Silica Powder	2	ġ.
Chalk Meal	1.5	
Vienna Lime	2	g.
Oxalic Acid	1	g.
Ammonia (28%)	10	cc.

Polish for Leather Furniture Paraffin Wax (50-52° C.) Ozokerite/Ceresin (58-60° C.) 5 g. 5 g. Wool Fat, Neutral 10 g. Becswax Carnauba Wax 10 g. 150 g. Turpentine Oil

Color similar to that of furniture. Pour at 40-45° C. into jar.

Furniture Polish

Formula No. 1		
Raw Linseed Oil	10	oz.
Spindle Oil	50	oz.
Stoddard Solvent	15	OZ.
Xylol	5	oz.
Soft Soap	1	oz.
Water	19	0 Z.
No. 2		
Paraffin Oil	20	oz.
Red Oil	5	oz.
C. M. C	•	

The above are mixed	vigorously	until
Water	70	OZ.
Gum Arabic	2	oz.
DOLL DOLL		UZ.

70 g.

completely emulsified.

No. 3	37
Carnauba Wax Montan Wax, Bleached	2 g. 6 g.
Beeswax	5 g.
Paraffin Wax (52-54° C.) Melt.	14 g.
Add:	
Linseed Oil (or Varnish)	8 g.
And (when temperatures 13	-45° C.∫

Turpentine

Liquid Furniture Polish		At same time prepare:	
Beeswax, Yellow	13 g.	c. Potassium Carbonate	5 g.
a. Ozokerite, Yellow	2 g.	Hard Soap	5 g.
b. Thinner (White Spirit)	75 cc.	Water, Hot	45 cc.
o. Alkali Solution (Water:		·	
monia (0.91) = 85:15	1) 10 ce	and pour in thin jet into a plu Keep temperature at $55-60^{\circ}$ C.	Stir con-
	-	tinuously, add a yellow dye, t	
Melt up a, add the warmed solution, then add c in thin j		into cans.	non pour
	et, stiffing	No. 2	
thoroughly.		Paraffin Scale	12 g.
Furniture Polish		Shellac Wax	5 g.
Formula No. 1		Carnauba Wax	4 g.
a. Paraffin Oil, Yellow	100 сс.	Ozokerite Ceresin (58-60° C.)	3 g.
Naphtha, Refined	50 cc.	Montan Wax, Bleached	4 g.
Tetralin, Dipentene	50 cc.	Turpentine Oil Substitute	72 cc.
Precipitated Chalk	25 g.	No. 3 (White)	
b. Lactic Acid (50%)	50 cc.		
Water	225 сс.	Carnauba Wax, Bleached	6 g.
Add b to a in thin, conti		Ozokerite, Refined Paraffin (50-52° C.)	4 g.
stir well.	naous jos,		20 g.
No. 2		Thinner (Turpentine Oils, Di-	1-0
Boiled Linseed Oil	10 lb.	pentene, Hydroterpene, Dec-	}70 g.
Raw Linseed Oil	12 lb.	aline, White Spirit)	,
Denatured Alcohol	2 lb.	No. 4 (White)	
Vinegar	12 lb.	Montan Wax, Double	
Turpentine	14 lb.	Bleached	12 g.
Petroleum Spirits	27 lb.	Montan Wax, Bleached	5 g.
or		Paraffin (50-52° C.)	6 g.
Raw Linseed Oil	2 gal.	Ozokerite, Refined	2 g.
Paint Drier	⅓ gal.	Thinner	75 g.
Vinegar	6 gal.	No. 5 (White)	
Furniture Finishers Po	lish	Montan Wax, Double Bleached	e ~
Turpentine	7 lb.	Montan Wax, Bleached	8 g.
Mineral Oil	7 lb.	Paraffin (50-52° C.)	3 g. 19 g.
Cedarwood Oil	2 oz.	Thinner	70 g.
Bassafras Oil	1 oz.	No. 6 (Yellow or Orange	
Rottenstone, Fine Powdered	4 oz.		•
		Carnauba Wax, Fat-Gray*	4 g.
Covering Polish for High-Glos	s Polished	Ozokerite, Yellow Paraffin (48-50° C.), Yellow	2 g. 24 g.
Furniture		Thinner	70 g.
Collodion Wool (Nitrocellu-	1	* Dye with 0.02% Sudan Yellow	
lose), Alcohol Soluble,	12 g.	2,0 mm 0.02,0 Eduar 10.00	u.
soaked in Butanol (2:1)	J	Tiquid Elean Delich	
Ethylene Glycol	6 g.	Liquid Floor Polish	
Toluene	12 g.	Melt:	
Tricresyl Phosphate	2 g.	Paraffin Wax (50-52° C.)	50 g.
Shellac (Free from Wax)	10 g.	Ceresin (58-60° C.)	10 g.
Alcohol (95–96%) or	E0 ~	Carnauba Wax	40 g.
Butanol Thinner (Alcohol)	58 g. optional	and dissolve:	
Zimmer (Medici)	opuoma	In summer, 7-9 parts in 93-	91 parts
		of turpentine.	
Floor Polish		In winter, 6-7 parts in 94-93	parts of
Formula No. 1		turpentine.	
Camauba Wax	15 g.	-	
	5 0	Deodorized Floor Polish	1
Rosing Pale	5 g.	Paraffin Wax (50-52° C.) 1	8 g.
Melt on Pale Melt on the bath, put out b. Turn cine Oil, or	fire. Add:	Carnauba Wax	5 g.
b. Turn dine Oil, or		Ceresin (58-60° C.)	2 g.
Spestitute	20 cc.	Rosin, Pale	4 g.
. —		•	

	POLISHES,	ABRASIVES 289
Stearin Potassium Carbonate Caustic Soda (38° Bé.) Water	1 g. 2 g. 0.5 cc. 66 cc.	No. 2 Spindle Oil, Refined (see above) 60 cc. Petroleum 27 cc.
Boil and stir until smooth.		Camphor Oil 3 cc.
Dyestuffs for Floor Po Yellow: Sudan Yellow RRN	lishe s	Spindle Oil, Refined (see above) 50 cc. Benzine 40 cc. Turpentine 5 cc.
Orange: Sudan Orange G, RR		Citronella Oil 5 cc.
Red: Sudan Red 5B		Mop Oil Polishes
Brown: Sudan Brown B, 3B, RRN		Above given formulae, but adding Waxes (as Montan Wax, Bleached, or Paraffin Scale
Reddish Brown: Sudan Brown 3B Sudan Red 5B	66% 34%	Wax) 2-3 g. Dye with Sudan Dyes 0.02 g.
Chocolate Brown: Sudan Brown 3B Sudan Red 5B Sudan Black BT	60% 30% 10%	Basic Dyes 0.06 g.
Other Oil Coloring Ba Yellow: Leather Yellow—Fat Dye		Water "Soluble" Floor Oil Spindle Oil, 5E (20° C.) 40 cc. Tallöl, Crude 20 cc.
Orange: Leather Yellow—Fat Dye Red Fat Dye Red:	66% 34%	Mix, warm to 70° C., add in thin jet: Caustic Soda, 38° Bé. 8 cc. Boil to saponify, add again Spindle Oil (as ahove) 27 cc.
Red Fat Dye Brown: Brown Fat Dye		Boil shortly, add boiling Water (to thin the alkali) 5 cc.
Reddish Brown: Brown Fat Dye Red Fat Dye	66% 34%	Use: 1 part oil in 6-10 parts water. Yellow Floor Wax
Chocolate Brown: Brown Fat Dye Red Fat Dye Ceres Black Lapieces Pigments:	60% 30% 10%	Formula No. 1 No. 2 No. 3 Paraffin Wax 16000 16000 16000 g. Carnauba Wax 3000 3000 2500 g. Beeswax, Yellow 1000 2000 1500 g. Turpentine 46000 25000 30000 cc.
Red: Iron Oxide Red Brown: Iron Oxide Brown		('Yellow 1435'' (Dye) 20 20 20 g. Amyl Acetate — 100 cc.
Floor Oils Spindle Oil, Pale, Viscosit 2.5-5E (20° C.), Ignitio Point 160-200° C. Olein	95 cc. 5 cc.	Dance Floor Wax Formula No. 1 Melt Paraffin Scale
Mop (Floor) Oils Formula No. 1	ı	G. (Yellow, 50-52° C.) 12 (1) Dye, Yellow or Red, Oil Soluble 25-56 g.
Spindle Oil, Refined, sp. g. == 0.850; 1.8-2.5E (20° C.) Benzine	70 cc. 25 cc.	Ochre, Yellow 8 g. Mix a and b ₄ thoroughly, coal, pulverize.
Balm-Turpentine Oil, Hydro terpene, Wood-Turpentin Oil or Refined Pine Oil	0- 10 5 cc.	Melt Paraffin Wax (50-52° C.) 80 g. Carnauba Wax, Refined 20 g.

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Sudan Yellow to suit Sudan Red Melt together, cool, pulverize.	Emery Polishing Paste Emery, Powdered 45 g. Aluminum, Powdered 4 g. Wax Paste Polish 24 g.
Linoleum Wax The following waxes are suitable for preservation of linoleum. The clear wax is also suitable as a floor wax or as a polish.	Wax Polish U. S. Patent 1,979,787 Carnauba Wax 9 1b.
Clear Wax Carnauba Wax 1 lb. 6 oz. Cerosin Wax 1 lb. 6 oz. Petroleum Spirits 8 lb.	Light Petroleum Oil 41 lb. Water 49.5 lb. Starch 0.5 lb.
Melt the two waxes together and stir in the petroleum spirits. The wax should then be ground. Red Wax	Wood Button Polish Turpentine 120 cc. Wax, White 120 g. Melt.
Carnauba Wax 1½ lb. Coresin Wax 1½ lb. Venetian Red, Dry ½ lb. Petroleum Spirits 6½ lb.	Add Alcohol 50 cc. with stirring. Axe or Hummer Handle Wax
Red Stain for Linoleum Venetian Red, in Oil 1½ lb. Boiled Linseed Oil 3 pt. Amyl Acetate 4½ pt.	White Beeswax 5½ lb. White Rosin ½ lb. White Lead 4 lb. Damar Varnish ½ lb.
Wax Polishes U. S. Patent 2,010,297 Formula No. 1 No. 2 Carnauba Wax 25 g. 2.75 g.	Melt the beeswax; crush, melt and stir in the rosin; add white lead while stir- ring, and finally pour in the damar var- nish. While still in a liquid state, this material is poured into small paper bags
Ceresin Wax 28 g. 3.08 g.	which serve as molds.
The four waxes should be melted to- gether at about 200° F., or somewhat higher, and the calcium stearate then dis- solved in the molten wax with gentle agi- tation. When the melt becomes clear, about half of the solvent is added. The	The finished product looks like bees- wax, but is lighter in color. The rosin and paraffin are melted and mixed and allowed to cool somewhat before stirring in the white lead and linseed oil—this to prevent foaming.

systallization occurs around 100-110° F.

The vigorous agitation is further continued until the batch reaches a temperature of 90-95° F., whereupon the other half of the solvent is slowly added in connection with gentle agitation. The product may then be packaged.

Wax Paste Polish 28 g. 6 g. 3 g. 4 g. Paraffin Ozokerite
Carnauba Wax, N.C. No. 3
Recewax, Yellow
Carpentine

solution is then cooled, to as low a temperature as 135-140° F. and vigorously agitated as by means of high speed stirrers, with the cooling continued until

60 cc.

Liquid Ski "Waxes" Formula No. 1

90 g. Shellac Sandarac 10 ğ. Alcohol 200 g.

Use solution to spread over the lower surface of the ski, from the top down, to about 10 cm. below the straps. Dry, and repeat spreading. For low temperatures, when snow has too much friction, add 1-2% Castor Oil.

No. 2

Carnauba Wax Montan Wax Linseed Oil Varnish

. t	OLISHES,	ABRASIVES	291
No. 3		No. 6	
Montan Wax, Refined	15 g.	Montan Wax, Crude	100
Ceresin	3 g.	Paraffin	120 g.
Turpentine Oil Substitute	82 g.	Wool Fat	30 g.
-	0	Seal Train Oil	20 g. 15 g.
No. 4	••	Tallow, Hard	10 g.
Colophony	30 g.	Rosin	5 g.
Ceresin	25 g.	Wood Tar	3 g.
Tallow	55 g.	No. 7	. B.
No. 5	10	Paraffin	1 g.
Talc	10 g.	Tallow	1.5 g.
Palm Oil	14 g.	Rosin	2.5 g.
Ceresin Paraffin	16 g.	Ozokerito	15 g.
No. 6	60 g.	No. 8	•
Tallow	105	Wool Fat	10 g.
	125 g.	Ceresin	90 g.
Colophony Montan Wax	275 g.	-	
	200 g.		
Turpentine Oil	200 g.	Ski Wax	
No. 7		Formula No. 1	
Rice Starch	40 g	1	
Tallow	125 g.	Montan Wax, Crude	18 g.
Larch Turpentine	260 g.	Paraffin Wax	60 g.
Yellow Wax	500 g.	Ozokerite	4 g.
No. 8		Wool Fat	6 g.
		Colophony	12 g.
(Sohm's Ski Wax)		Melt together and add turp	entine oil
Ozokerite	55 g.	to desired consistency.	
Tallow	15 g.	1	
Rosin	30 g.	No. 2	
All these waxes may be the	inned with	Ascension Wax:	
turpentine oil to desired fluids	ity.	Ceresin	10 g.
_		Paraffin Wax	20 g.
1-1-		Wool Fat	28 g.
Norwegian Klister (Ski)	Waxes	Colophony	15 g.
•	TI WALCO	Montan Wax	27 g.
Formula No. 1		Melt together and add turn	
Rice Starch	40 g.	desired consistency.	ALIEN CO
Tallow	125 g.	devired consistency.	
Larch Turpentine	260 g.	No. 3	
Yellow Wax	500 g.	Gliding Wax:	
No. 2		Paraffin Wax	20 -
Paraffin (40-42 C.)	60 g.	Ceresin	60 g.
Colophony	12 g.	Tallow	16 g.
Wool Fat	6 g.	ľ	14 g.
Carnauba Wax	4 g.	Melt together and add turp	entine to
Montan Wax	80 g.	suit.	
No. 3	о в.	No. 4	
	EE	Gliding Wax:	
Ozokerite Colophony	55 g. 35 g.	Black Ozokerito	55 g.
	10 g.	Rosin	30 g.
Spindle Oil, Refined	10 g.	Tallow	15 g.
No. 4		Melt together and add turn	entine to
Paraffin	70 g.	suit.	
Colophony	15 g.	No. 5	
Wool Fat	10 g.	Paraffin Wax	30 ~
Carnauba Wax	5 g.	Montan Wax, Bleached	30 g. 80 g.
Montan Wax	15 g.	Colophony ·	20 g.
No. 5		Japan Wax	20 g.
Ozokerite	5 g.	Wood Tar Oil	20 g. 10 cc.
Colophony	4 g.	Turpentine Oil	10 cc.
Train or Spindle Oil	1.5 g.	Yellow Dyestuff enough	
or phindip on	v 9.	chough	M COLUL

				<u>.</u>
No. 6	,	High-Luster Polish for	Shoes	_
Wax Polish, White:		Formula No. 1		
Paraffin Wax	16 g.	Carnauba Wax, Yellow	500	~
Carnauha Wax, Light	3 g.	Carnauba Wax Residue	500	
Beeswax, White	1 g.	a. Montan Wax, Bleached	500	Ç.
Turpentine	46 cc.	Paraffine (50-52° C.)	200	<u>چ</u> .
No. 7		Colophony	150	ž.
Wax Polish, Liquid:		Water	8500 c	
Paraffin Wax	50 g.	b. Potash, Caustic	300 g	
Ozokerite	5 g.	Olive Oil Soap	100 g	
Carnauba Wax	100 g.	c. Turpentine Oil or	6	•
Turpentine Oil	750 cc.	Substitute	1500 c	e.
Benzoline	94 cc.			
Camphor Oil	2 g. 3 cc.	Melt up a, saponify with b		
Amyl Acetate	3 cc.	cool, and add c, shortly before	solidi:	ned.
No. 8		No. 2		
For Gliding:		Montan Wax, Crude	6 g	7.
Paraffin (50-52° C.)	60 g.	Carnauba Wax	3 0	7.
Ceresin (60° C.)	16 g.	Ozokerite (58-60° C.)	2 8	ζ.
Tallow or Palm Oil	14 g.	Candelilla or Shellac Wax	3 g	ζ.
Talcum No. 9	10 g.	Paraffin Scales (50-52° C.)	14 g	ζ.
For Climbing:		Nigrosine Base	3 g	ζ.
	* 0 -	Turpentine 2	20-30 c	æ.
Paraffin (40–42° C.) Rosin	50 g. 20 g.			
Wool Fat	15 g.			
Wood Tar -	15 g.	Shoe Polish Paste		
No. 10	B.	Carnauba Wax, Fat-Gray	6 g	z.
Climbing and Sliding Ski W	ov.	Montan Wax, Bleached	7 o	7.
		Paraffin (50-52° C.)	11 2	ž.
Paraffin Montan Wax, Crude	40 g.	Ozokerite	2 g 2 g	ζ.
Wool Fat, Neutral	15 g. 15 g.	Dyestuff	2 g	ζ.
Rosin	10 g.	Thinner (Turpentine Oil, or		
Mineral Oil	15 g.	substitute or a Mixture of Both)	72 c	
Wood Tar	5 g.	1000)	12 0	٠.
No. 11	•			
Climbing Wax:		Shoe Polish		
Montan Wax, Crude	17 g.			
Wool Fat, Neutral	18 g.	British Patent 395,53	8	
Paraffin	10 g.	Paraffin 💏	14 g	ز ٠
Rosin	28 g.	Ozokerite	3 g	۲.
Ozokerito Mineral Oil	25 g.	Carnauba Wax	3 g	5 -
Wood Tar	5 g. 2 g.	Melt 80-90° C.		
	- g.	Turpentine Oil	38 c	c.
		Stir now with		
Ski Finishes		Water (boiling)	38 c	c.
For running on wet snow.		Sodium-Sulphonate of Glycol		
Mix:		Mono-Oleate,	1 g	5•
Pine Tar	25 g.			
Copal Lacquer Venice Turpentine	25 g. 50 g.	Dyeing Shoe Polish, Lie	quid	
This mixture is boiled in		Carnauba Wax, Fat-Gray	2 g	۲.
ning side of the ski with a	blowtorch.	Montan Wax, Bleached	2 g	
Before using the ski rub in	a thin coat-	Paraffin (50-52° C.)	4 g	
ing of Venice turpentine.		Ozokerite, Refined	1 g	; -
For running on very cold		Dyestuff Thinner (Tunnentine Oil on	1.5 g	:-
in a good coating of Pine tar using heat ski and rub on a		Thinner (Turpentine Oil, or Substitute, or Mixture of	,	
aceti.	omo shorm.	Both)	89.5 c	c_
		/		

Sporting Shoe Dressings, Paste	Paraffin (50-52° C.) 9 g.
Formula No. 1	Black Dye, Oil-Soluble 3 g.
Shoe Paste, Black	Thinner (see above) 60 cc.
_ •	Spindle Oil, Refined 15 cc.
Carnauba Wax, Gray 7 g. Montan Wax, Crude 7 g.	
Paraffin (50-52° C.) 12 g.	No. 3
Black Dye, Oil-Soluble * 3 g.	Carnaula or Shellac Wax 8 g.
Thinner (Turpentine Oil, or	Montan Wax, Crude 8 g.
Substitute, or a Mixture of	Paraffin (50-52° C.) 10 g.
Both) 51 cc.	Black Dye, Oil Soluble * 3 g.
Vaseline Oil 20 cc.	Thinner (see above) 51 cc.
No. 2	Spindle Oil, Refined 10 cc.
Carnauba Wax 5 g.	Sardine Tiain Oil 10 cc.
Montan Wax, Crude 8 g.	
	TO 11 1 T 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
.4-74	Polishes, Liquid
Fo	rmula, No. 1 No. 2 No. 3
Carnauba Wax, N.C.	3 g. 3 g. 4 g.
Montan Wax, Crude Paraffin (50-52° C.)	2 g. 2.5 g. 2 g.
Paraffin (50-52° C.)	3 g. 2.5 g. 2 g.
Black Dye,* Oil Soluble	3 g. 3 g. 3 g.
Thinners (see above)	75 cc. 72 cc. 70 cc.
Spindle Oil, Refined	14 cc. — 19 cc.
Vaseline Oil	- 11 cc - 6 cc
Sardine Train Oil	- 6 cc
* Black Dyes	b. Montan Wax, Crude 7 g.
Nigrosine Base 51017 Nigrosine Base 4322	Japan Wax 2.5 g.
Nigrosine Base 4322 Nigrosine Base LJF	Carnauba Wax, Gray 4 g.
Nigrosine Base SRN	Beeswax 2.5 g.
Nigrosine Base SR Nigrosine Base C	Paraffin (50–52° C.) 2 g. Od Soluble Black 2.5 g.
	0.2
How to dissolve the Black Dye:	Pour b molten into hot a. To the
a. Olein 1 g.	homogeneous (cooled) mass add while
Montan Wax, Raw 1 g.	stirring
Nigrosine Base 1 g.	c. Turpentine 25 cc.
Warm together and stir.	
b. Stearin 2 g.	Notes on Cleaning White Shoes
Nigrosine Base 1 g.	Important note-all cleaners should be
Trigiositic Daso - 8.	applied sparingly. It is best to place the
Black Shoe Polish	shoes to be cleaned on the shoe trees and
Carnauba Wan 6 g.	with a dry cloth remove surface dust or
Montan Wax, Crude 5 g.	dirt. Do not clean white shoes while on
Soft Ozokerite (58-60° C.) 1 g.	the feet.
Nigrosine Base 3 g.	Apply the cleaner sparingly to a clean
Paraffin (58-60° C.) 14 g.	white cloth, preferably toweling, and first
Turpentine 71 cc.	clean the dirtiest spot, then go all over
***************************************	the shoe, using sufficient pressure to re-
Powder Glaze for Shoes	move all spots and stains. Avoid satu-
Shellac 18 g.	rating the leather but apply evenly over the entire area to be cleaned.
Borax • 71/2 g.	Permit shoes to dry thoroughly. Next
Water 75 g.	rub the shoe briskly with a clean dry
Dissolve and men evaporate water	cloth, removing all white particles of
until dry and then pulverize.	powder and until the original sheen is re-
	stored.
Shoe Cream for Collapsible Tubes	In the case of white buck or suede
a. Water 🖢 52 cc.	shoes, a fine bristle brush will more easily
Nigrosine 1 g.	remove excess powder and raise the nap
Potassium Carbonate 0.5 g.	of the leather.
Hard Soap 0.75 g.	Do not use soap and water on elk shoes.
Boil.	Beware of a cleaner with so much alkali

that repeated usage will remove, the finish. This generally results in the hardening of the clk leather so that it cracks or shrinks.

White Shoe Polishing Stick

Carnauba Wax, Flora	4 lb.
Stearic Acid	4 lb.
Paraffin Wax	17 lb.
Montan Wax, Bleached	16 lb.
China Clay	9 lb.
Titanium Dioxide	1 lb.

White Shoe Dressing

Titanox A	10.5 oz.
Titanox B	20.75 oz.
White Soap	3 oz.
White Dextrin	3 oz.
Ammonia	1.25 oz.
Water	48.40 oz.
Carbon Tetrachloride	13.25 oz.
Moldex or Other	
Preservative	10 oz.

Shoe White (Water Type)

This cleaner for white canvas and leather shoes cleans and whiteus at the same time and leaves a coating which does not dust or rub off.

roce not duet of the on-		
Lithopone	28	oz.
Asbestine	4	oz.
Gum Arabic	7.5	oz.
Gum Tragacanth	0.3	oz.
Benzoate of Soda or Moldex	0.5	oz.
Ultramarine sufficient to	o whi	ten
Perfume		

sufficient to give pleasant odor Water 59.7 oz.

If better hiding power is desired titanium dioxide pure or titanium dioxide with a barium or calcium base may be used; as well as pure zinc sulphide. The asbestine is added to prevent the pigment from packing hard on long standing. The tragacanth gives added body or viscosity, and inhibits much of the pigment from settling, a mere inversion of the bottle being adequate to bring same back into suspension.

Shoe White (Waterproof Type)

This composition leaves a coating which is waterproof and does not dust off. It is preferred to the water type for leather shoes particularly the glazed type.

Lithopone Asbestine		28 oz. 4 oz.
Ultramarine sufficient Ester Gum, Pale	to	whiten 5 oz.

62 oz. 1 oz.

sufficient to mask petroleum odor
The solvent naphtha should be a petroleum fraction boiling between 200° and
300° F. The aluminum stearate is dissolved in same to increase the viscosity
and inhibit settling of the pigments.
The ester gum is then added and stirred
or heated until solution is complete. The
perfume and pigments are then added.

White Shoe Cleaner

THE SHOO CICUICI	
a. Titanox C b. Diglycol Laurate	30 g.
o. Pigiy of Daulate	6 cc.
a molene	10 cc.
Tolhol	12 cc.

Mix a and b thoroughly.

	Bright	Drying Carnauba		
*		Emulsion	60	cc.
	Water		20	cc.

Add c to ab in 4 equal portions, shaking or stirring during and after each addition.

d. Trichloroethylene 40 cc.
Add slowly with stirring.

White Shoe Dressing

Titanium White	60 g.
Diglycol Oleate	12 g.
Naphtha	20 g.

Stir the above together and while stirring vigorously add slowly

Carnauba Wax Emulsion

(10% Wax)	80 g.
hen stir in vigorously	
(Daiable access to the control of th	40 400

Trichloroethylene 60-100 g.

Polishing Cloths

Prepare powder mixtures:

Formula No. 1	
Calcium Carbonate Kieselguhr Caput Mortuum	70 g 25 g 5 g
No. 2	
Magnesia, Calcined English Red	20 g 40 g
Vienna Lime	40 g

No. 3	
Calcium Carbonate	40 g.
Bolus	20 g.
Vienna Lime	20 g.
Infusorial Earth	10 g.
Magnesia Usta	5 g.

One hundred and fifty grams of these mixtures are stirred into 1000 cc. of

ater, impregnate the cloths in this susension. Press. Dry (40-50° C.). Fix ith a bath of 100 g. hard soap in 1000 c. water. Press and dry again.

Cleansing and Polishing Compositions British Patent 425,323

A cleansing and polishing liquid which caves a thin film on the leather, wood, metal, or other article treated, is composed of a hard wax polishing composition, alkali, water, a solvent of oil and fat, carbon tetrachloride shellae, and bornyl acetate. For example, 3 lb. of shellae wax, 3 lb. of montan wax, 3 lb. of carnauba wax, 2 lb. of parafiin wax, 1 lb. of japan wax, 1 lb. of acetone varish, 1 lb. of nitrocellulose varnish, 1 lb. of cellulose varnish, 3 lb. of potash or soda, 20 lb. of water, 1 lb. of castor oil, 5 lb. of white spirit, 40 lb. of turpentine substitute, 20 lb. of carbon tetrachloride, 1 lb. of shellae, and 1 lb. of bornyl acetate are mixed together.

Pore Filler for Polish Bases German Patent 607,521

Carnauba Wax	5 g.
Punice Powder	100 g.
Sandarac	100 g.
Castor Oil, Blown	10 g.
Shellac Wax	10 g.
35.34	and and au

Melt up while stirring, cool, and pulverize. The "Pore Filler" is then ap-

plied as usual by rubbing it in on the wood surface together with the polishing liquid.

Abrasive Wheel Formula No. 1 British Patent 411,846

One hundred parts abrasive grains are coated with 1 part of a resin solvent, e.g., di-butylphthalnet, and 6-20 parts of finely divided glycerol-phthalic anhydride reaction product are added, the mixture is warmed to 350° F. to make it plastic and passed several times between rollers, covered with a thin film of linseed oil and maintained at 150° F., and, after final sheeting, articles are cut out and hardened for 48 hours at 350° F.

No. 2

British Patent. 434,402

251111011 2 411141	-
Diamond Dust	26 oz.
Graphite	50 oz.
Charcoal	50 oz.
Red Iron Oxide	75 oz.
Phenol Formaldehyde Resin	
sufficient	to bond

Hardness Scale for Abrasives

A scale of hardness based on the lapping method is as follows: bort 10, ballas 9.99, carbonado 9.82, boron carbide 9.32, black silicon carbide 9.15, corundum 9.00.

Fireworks (Pyrotechnics)

The greatest care should be exercised in making fireworks. Curelessness and impurities produce most accidents. Do not mix large amounts of ingredients and do not permit the introduction of dirt, dust or other foreign matter. Do not mix near your stock of raw or finished material. Make sure that all utensils are cleaned directly before use. Slight friction, even that produced by sifting may cause an explosion or fire. All packing or ramming should be done gently and without scratching as the latter may start a reaction just as well as a shock. Do not allow matches or open flames

Do not allow matches or open flames in the mixing room. Wear rubber soled shoes. Keep the air moist enough to prevent static sparks from being generated

by moving bodies.

All chemicals used should be of best quality and bought from a reliable house in original packages. These should be kept air-tight. For mixing small quantities round brass wire sieves (No. 16-26) are used. In plain mixings the coal is weighed first and put into bottom of a wooden tub; the sieve is put on top and the sulphur and saltpeter sifted through it. Then with bare arms mix the powder in the tub thoroughly. Place sieve on another tub and sift from first tub a scoopful at a time. Mix with hands again and sift back again into first tub.

"In "colored?" mixings each ingredient should be sifted separately the first time except the shellac, coal, etc., which is put in bottom of tub. Never throw the chlorate on the sieve with dextrin or other organic material. Beware of hitting the sieve with finger nails or metallic objects.

Sparklers

Fo	rmula No. 1	No. 2
Lampblack		— lb.
Powdered Charcos		25 lb.
Steel Filings		50 lb.
Aluminum Powde	r 15	lb.
Gum Arabic	6	5 lb.
Saltpeter	5	15 lb.
Sulphur	2	6 lb.

The gum arabic is worked up with water into the consistency of mucilage, the other items except the steel filings are stirred in. The steel filing lightly coated with paraffin is finally added. Then work the mixture up to the consistency of porridge.

Pin Wheels

Formula	No. 1	No. 2	No. 3
Meal Powder		10	8 lb.
Fine Grain Powd	er 8	5	8 lb.
Aluminum			3 lb.
Saltpeter	14	4	16 lb.
Steel Filings	6	6	— lb.
Sulphur	4	1	3 lb.
Charcoal	3	1	8 lh.

Pyrotechnic Fountains

Tyrotechnic Tountains		
Meal Powder	5	lb.
Granular Saltpeter	3	lb.
Sulphur	1	lb.
Coarse Charcoal	1	lb.
FF Rifle Powder	%	lb.

Flower Pots

Saltpeter	10 lb.
Sulphur	6 lb.
Lampblack	3 lb.
FFF Rifle Powder	6 lb.

Gerbs

	Formula.	No. 1	No. 2
Meal Powder		6	4 lb.
Saltpeter		2	— lb.
Sulphur		1	lb.
Charcoal		1	1 lb.
Steel Filings		1	2 lb.

Serpents or "Nigger" Chasers

	Formula	No. 1	No	. 2
Meal Powder	#	3	3	lb.
Saltpeter		2	5	lb.
Sulphur		1	1	lb.
Mixed Coal		11/2	8/4	lb.
FFF Grain P	owder	4	3	lb.

Saltpeter Ammonium Bichromate Dextrin	-	1 lb. 2 lb. 1 lb.
Table Rocke	t	
Formula:	No. 1	No. 2
Saltpeter Meal Powder Charcoal Sulphur Steel Filings	2 2	5 lb. 12 lb. 3 lb. 3 lb. — lb.
Roman Candle	e s	
Powdered Saltpeter Fine Powdered Charcoal Flowers of Sulphur Dextrin Water	1	8 lb. 1 lb. 5 lb. 1 lb. 1 gal.

Rocket and Candle Match

evenly dampened.

After all the ingredients are well mixed and sifted 3 times, add the water and mix again until the whole lot is

Into a small tub put about a gal. of starch, well boiled, and stir into it about 5 lb. of a thoroughly mixed composition made of

Saltpeter	16 lb.	
Fine Charcoal	5 lb.	
Sulphur	21/2 lb.	
Soak in this cotton w	rick of about	5

strands until nearly all the composition is absorbed but about 1/2 in. should still cover the cotton in the tub.

Cascades	l	
Formula.	No. 1	No. 2
Granulated Saltpeter	18	16 lb.
Mixed Charcoal	4	4 lb.
Sulphur	3	3 lb.
Iron Borings	6	7 lb.
Smoke Po	ot	
Strontium Nitrate		10 lb.
Sulphur		6 lb.
Whiting (Chalk)		4 lb.
Fine Charcoal		¾ lb.
Dextrin		% lb.
or		
Saltpeter		4 lb.
Lampblack		1 lb.
Charcoal		1 lb.
Red Arsenic		1 lb.
Rosin		1 lh

Gold and Silver Rain

(Cut Stars)

Formula.	No. 1	No, 2	No. 3
Meal Powder	16		4 lb.
Saltpeter	10	1	1 lb.
Sulphur	10	1	— lb.
Fine Charcoal	4	1	2 lb.
Lampblack	2		— lb.
Red Arsenic	1		lb.
Shellac	1	_	lb,
Dextrin	1		lb.
Lead Nitrate		3	- lb.

Japanese St	ars	
Formula	No. 1	No. 2
Lampblack	12	6 oz.
Potassium Chlorate	8	4 oz.
Saltpeter	1	- oz.
Water	18	9 oz.
Alcohol	4	2 oz.
Dextrin	1	oz.
Gum Arabic	_	1/2 oz.
201 41 2 4 10 10 2 10	144	

Mix the dextrin and saltpeter together and add sufficient water to make a gummy liquid. Boil the balance of the water and add the potassium chlorate to it. Put the lampblack in a large pan and pour the alcohol over it working it in as well as possible. Then add the potascum blasters in the het water and the sum chlorate in the hot water and stir with stick until cool enough for the hands and lastly add the dextrin and saltpeter.

In Formula No. 2 the potash and lampblack are sifted together several times; add alcohol; then water in which gum has been dissolved and proceed as in Formula No. 1.

White Stars

White Sta		
Formula	No. 1	No. 2
Saltpeter Sulphur Red Arsenic Dextrin Black Antimony Red Lead Shellac	50 15 15 3 —	54 lb. 15 lb. 9 lb. 3 lb. 15 lb. 6 lb. 1 lb.
Red Stars		No. 2
Tormula .	110. 1	110. 2

Formula	No. 1	No	. 2
Potassium Chlorate	6	24	lb.
Shellac or Red Gum	1	3	lb.
Fine Charcoal	2	4	lb.
Strontium Carbonate	_	4	lb.
Strontium Nitrate	в		lb.
Dextrin	1/2	11/6	lb.

Blue Stars Potassium Chlorate 24 lb. Paris Green 9 lb. Barium Nitrate 8 lb. Shellac 5 lb.	Each ingredient should be sifted sepa- rately and then mixed in a tub with the fingers, preferably gloved, being careful not to scratch the bottom of tub with the nails.
Dextrin 1½ lb. Chinese Fire Crackers	Japanese or Cap Torpedoes Formula No. 1
Formula No. 1 No. 2 Saltpeter 50 45 lb. Sulphur 25 18 lb. Charcoal 25 25 lb. Potassium Chlorate 8 lb. Sand - 4 lb. Flash Crackers Formula No. 1 No. 2 No. 3 Saltpeter 50 lb. Sulphur 30 25 30 lb. Aluminum Powder,	Potassium Chlorate 5 oz. Sulphur 4 oz. Chalk No. 2 Amorphous Phosphorus 2 oz. Sift separately the ingredients of No. 1, mix thoroughly and moisten in a bowl with water until of the consistency of porridge. In another bowl moisten the 2 oz. of amorphous phosphorus, to the same consistency. Then stir the phosphorus into the bowl containing the other
Fine 20 25 40 lb. Potassium Chlorate — 50 30 lb. Cannon Cracker Composition Formula No. 1 No. 2 No. 3	ingredients with a spoon. White Fire Formula No. 1 No. 2 No. 3 No. 4
Potassium Chlorate 60	Saltpeter 3 12 8 7 lb. Sulphur 1 2 2 2 lb. Metallic Antimony 1 - - lb. Sulphide of Antimony 1 1 - - lb. Realgar - - 1 14/2 lb.
Red Formula Nitrate of Strontia. Potassium Chlorate Shellac Sheel-lac or Kauri Gum Charcoal Dextrin Fine Sawdust Rosin Lampblack	No. 1 No. 2 No. 3 No. 4 No. 5
Blue Formula, 1 Chlorate of Potash Paris Green Stearin Shellac Barium Nitrate	No. 1 No. 2 No. 3 No. 4 6 8 8 12 lb. 4 6 6 8 lb. 1 1 1 1 2 lb ½ ½ - lb.
Calomel Sal Ammoniac In order to make tableau fires more bulky, one to two parts of fine sawdust may be mixed with any of the above formulas without materially affecting the	that paris green is very poisonous and a

PYROTECHNICS	
Green Fire	Railway Fuses
Formula No. 1 No. 2 No. 3	Formula No. 1 No. 2 No. 3 No. 4
	Strontium Nitrate 48 16 18 16 lb.
Barium Nitrate 8 9 4 lb. Potassium Chlorate 4 3 2 lb.	Saltpeter 12 4 7 4 lb.
Shellac — 1 1½ lb.	Sulphur 5 2 2 5 lb.
Shed-lac (Shellac	Fine Charcoal 4 1 1/2 1 lb.
Substitute) 2 — lb.	Fine Charcoal 4 1 1/2 1 lb. Red Sheel-lac 10 3 2 — lb.
Dextrin — 1/16 — lb.	Dextrin $ \frac{1}{2}$ $ \frac{1}{2}$ $ \frac{1}{2}$ $ \frac{1}{2}$
Fine Sawdust - 1/2 - lb.	Accompany of the same of the same
Sal Ammoniac 1 — lb.	Ship Distress Signals
	Potassium Chlorate 5 lb
Yellow Fire	Strontium Carbonate 11/2 lb.
Barium Nitrate 36 lb.	Shellac 1 lb.
Sodium Oxalate 6 lb.	Dextrin 11/2 lb.
Sulphur 3 lb.	
Sheel-lac 5 lb.	Miracle Candles
	a. Iron, Powdered 25 g.
Red Lances	b. Barium Nitrate 52 g.
Formula No. 1 No. 2	c. Aluminum Powder 8 g.
Potassium Chlorate 16 16 lb.	d. Starch, Wheat 15 g.
Strontium Nitrate 3 - lb.	Right size of the iron grains is most
Strontium Carbonate — 3 lb.	important, b and c should be finely
Shellac 3 2 lb.	powdered.
Lampblack 1/8 1 lb.	Should be produced in summer for quicker and more economical drying.
	Mixture must be perfect, pack in air-
Green Lances	tight drums,
Formula No. 1 No. 2 No. 3 No. 4	Put into an enameled container (best
Potassium Chlorate 7 16 16 - lb.	way using 1 kg. mass), make a little hole
Barium Nitrate 7 4 6 — lb.	in the center of the powder, pour in it
Barium Chlorate 6 lb.	the least possible amount of boiling
Shellac 2 4 3 1 lb.	water (100 g. for 1 kg. powder), and stir
Calomel — 3 3 2 lb. Lampblack — ½ — lb. Dextrin — 1 — lb.	the whole thoroughly. The right point of pastification and right amount of
Lampblack — 1/8 — — lb.	water is reached when the paste is not
	too friable or too sticky and forms a con-
Pierie Acid — 1 1 lb.	crete non-sticky mass.
	This mass is put on wires (2 g. per
White Lances	wire), and dried.
Formula No. 1 No. 2 No. 3 No. 4	
Saltpeter 9 14 5 8 lb.	Orange Smoke
Sulphur 1 4 2 2 lb.	U. S. Patent 1,975,785
Antimony Sulphide 2 — — — lb.	1
Antimony Metal	A pyrotechnic composition for pro- ducing orange smoke, comprises lead
Powder — 3 1 — lb, Meal Powder — 1 — lb.	ducing orange smoke, comprises lead
	peroxide 50 parts, potassium bichromate
Red Arsenic — — 1 lb.	35 parts, and magnesium 15 parts.
	P. G. L.
Magnesium Torches	Brown Smoke
a. Shellac 120 g.	U. S. Patent 1,975,099
Resin 120 g.	A pyrotechnic composition for pro-
Barium Nitrate, Dry 840 g.	ducing brown smoke, comprises copper
b. Magnesium Powder 25-40 g.	oxide 50 parts, lead peroxide 35 parts,
Mix the ground a with b, and fill into	and magnesium 15 parts.
zinc-tubes (thin walls) having a wooden	
handle, which closes the tube below.	Pyrotechnical Device

Parade Torches

Strontium Nitrate Potassium Chlorate Red Sheel-lac 40 lb. 8 lb. 7 lb.

Pyrotechnical Device U. S. Patent 1,936,221

A firework of the "sparkler" type consists of an iron rod coated at one end with a plastic mixture of barium nitrate

85, strontium carbonate 60, sodium aluminum fluoride 40, potassium chlorate 225, dextrin 30, and shellac 55 all parts by weight in which are embedded granules of magnesium-copper or magnesiumaluminum alloy.

Explosives

Formula No. 1 British Patent 408,260

Explosives consist of alpha-trinitrotoluene 10-30, o-nitrotoluene 5-10, ground coconut fiber or charcoal 1-5, o-nitrotoluene 5-10, paraffin or other suitable wax 3-6, aluminum, graded 50-mesh, 10-24, finely powdered aluminum 1-4, and barium nitrate, or other nitrate, 70-21 parts.

No. 2

British Patent 412,583

A nitrated mixture of glycerol and glycol 15, ammonium nitrate 8.5, sodium nitrate 12.0, plant fiber 6, sodium chloride 58 and ammonium orthophosphate 0.5%, has a density of 1.1 g. per cc. and gives a ballistic pendulum swing of 1.08 in., the volumetric power factor being 1.19.

No. 3

British Patent 435,588

Ammonium Nitrate QΛ lb. Aluminum Powder 61/2 lb. Manganese Dioxide Powder 31/2 lb.

Slow-Burning Explosives British Patent 423,040

Examples of slow-burning explosives are (1) potassium nitrate 75 or sodium nitrate 73, charcoal 15 or 17 and sulphur 10, (2) sodium nitrate 44, ammonium nitrate 34 and charcoal 22%. The explosive may be granular or compressed in pellets and may contain small quantities of cooling salts and boric acid or borates.

Explosive Priming Mixture British Patent 432,096

A suitable composition is the potassium salt 16, basic lead salt of trinitroresorcinol 15, barium nitrate 40, and antimony sulphite 29%.

Priming Charge Canadian Patent 348,291

A solution containing potassium ni-trate 30, barium nitrate 20, and water

100 parts is crystallized at 50° C. to give a double salt, which when used in priming charges leaves substantially no corrosive residues nor fused masses in the barrels of firearms; e.g., a priming charge consists of mercury fulminate 20-45, potassium barium nitrate 30-60, lead thiocyanate 10-40% by weight.

Priming Composition German Patent 614,712

The composition contains zirconium powder in addition to the usual constituents. Thus, the composition may contain zirconium powder 10, barium nitrate 40, mercuric fulminate 25 and antimony trisulphide 25%.

Flash Composition U. S. Patent 1,964,077

A suitable mixture contains perchlorate 20, potassium chlorate 39.5, silver nitrate 39.5, and nitrocotton 1.0%.

Flashlight Cartridges British Patent 419.658

A cartridge is charged with a powder mixture consisting of magnesium 700-900, sulphur 10-18, potassium permanganate or potassium chlorate 100-140, potassium nitrate 70-85, magnesium oxide 100-160 and charcoal 10-13 parts.

Black Powder Canadian Patent 348,641

The addition of 0.1-5.0% by weight of stearic acid retards the burning speed of black powder. E.g., a black blasting powder contains sodium nitrate 72.0, sulphur 10.0, charcoal 17.7 and stearic acid 0.3%.

Fuse Powder

French Patent 783,249

A powder of long combustion is made by dissolving niter 5, pulverized sulphur 4 and wood charcoal 3.5 parts in pure alcohol to form a thick mass which is well mixed and dried.

Gelatin Dynamite Canadian Patent 352,763

The following percentage compositions are specified:

Formula No. 1

Nitroglycerin Dinitrotoluene

Nitrocotton	1.3
Sodium Nitrate	36.1
Expanded Cereal Product	9
Starch	2.7
Chalk	0.9
No. 2	****
Nitroglycerin	60
Dinitrotoluene	3.5
Nitrocotton	2.3
Sodium Nitrate	2.2
Ammonium Nitrate	24
Expanded Cereal	6
Starch	1
Chalk	1
No. 3	
Nitroglycerin	30
Dinitrotoluene	2
Nitrocotton	0.7
Sodium Nitrate	44.8
Ammonium Chloride	15
Expanded Cereal Product	2
Starch	4,5
Chalk	1
No. 4	
Nitroglycerin	22
Dinitrotoluene	1.5
Nitrocotton	0.2
Sodium Nitrate	9
Ammonium Nitrate	60
Expanded Cereal Product	69
Chalk	0.4

Detonators French Patent 781,646

A composition which is fired directly by the passage of an electric current comprises a mixture of finely divided zirconium and a mtrophenol salt of lead, e.g., zirconium 70 and lead mononitroresorcinate 30 parts in sufficient amount of a 5% solution of nitrocellulose in amyl acetate to make a creamy mixture.

Percussion Detonator U. S. Patent 1.975,679

A percussion detonating composition consists of phosphorus sesquisulphide 30 g., gum arabic 115 g., magnesium carbonate 20 g., calcium carbonate 5 g., potassium chlorate 80 g., iron sesquioxide (red ochre) 40 g.

Waterproofing for Blasting Fuses (Non-Staining) U. S. Patent 1,968,907

Paraffin Wax

25-90 lb. Petrolatum Ester Gum 5-75 lb. 5-50 lb.

RUBBER, RESINS, WAXES, PLASTICS

Caoutchouc (Rubber) Synthetic Acetylene is absorbed by a mixture of 1000 g. Cuprous Chloride Ammonium Chloride 400 g. 100 g. Copper Hydrochloric Acid 30 g. Concentrate 425 g. Water at 40-50° C. The saturation is reached, when 50 g. of acetylene have been absorbed (3 hours). The mixture is kept at ordinary temperature during 24 hours, then distilled on an oil bath. The distillate contains 33% of

CH₂ = CH→ C = CH, and 67% of superior condensation products, among which has been found

 $CH_2 = CH - C = C - CH = CH_2$, and C_8H_8 .

In the same process, the yield in CH₂ = CH -- C ≡ CH

falls, when the period between saturation and distillation is increased to 140 hours. A 70% yield is obtained when running the absorption at 80° C., and collecting the gas of reaction into two receivers, the first chilled in ice, the second in carbon dioxide snow. The liquid in the

second receiver contains:

CH₂ = CH - C = CH

CH = CH

CH₃ - CHO

The chloroprene is obtained with an 80% yield, agitating

 $CH_2 = CH - C = CH$ with Hydrochloric Acid,

Concentrate 70 g.
Cuprous Chloride 10 g.
Ammonium Chloride 4 g.
for 3 hours at room temperature.

Rubber Master-Batch

U. S. Patent 1,942,853

Substantially unmasticated crude rubber (1 lb.) is superficially with 14-3 lb. of a softener, &g., mineral oil, so that the latter is absorbed. This procedure obviates the difficulties of incor-

poration of liquid softeners in the usual manner, and the soft, non-tacky product is very easily mixed with other compounding ingredients.

Porous, Fibrous Rubber Compositions British Patent 409,294

A porous, non-waterproof, fibrous, feltlike material is prepared by admixture of rubber with finely comminuted (not powder) fibers of wool and hair in proportions of not more than 50% rubber and not less than 50% fibers together with an amount of non-liquid expanding agent, e.g., ammonium carbonate, sodium carbonate, sodium potassium carbonate, sodium bicarbonate, ammonium bicarbonate that will expand the mass 2-6 times. Vulcanization and coloring agents and softeners may be added. In an example sulphur 7.5, zinc oxide 6, ferric oxide 2, stenric acid 4, ultra-accelerator 1 oz. and comminuted wool 22.5 lb. are added to 15 lb. softened rubber. When cool ammonium bicarbonate is added and the product calendered into sheets.

Rubber Fibers German Patent 614,615

Rubber fibers are formed by introducing a coagulating agent through nozzles into rubber latex. Thus, a 60% solution of acetic acid is fed into a rubber latex mixture of rubber 92.5, sulphur 2.5, zinc oxide 2.5, anti-oxidation agent 1.0, accelerator 0.5 and ammonium cleate 1%, through 0.42 mm. nozzles, the fiber being removed at 600-760 cc. per minute.

Chlorinated Rubber British Patent 410,249

A solution of unvulcanized (artificial or reclaimed) rubber, gutta-percha or balata, with or without factice, admixed with 5-20% uncombined sulphur is chlorinated to yield a thermoplastic mass suitable for the manufacture of films, varnishes or moldable compositions,

the chlorination being continued until the gel which forms is entirely redissolved. Metallic halides, oils, turpentine, chlorinated naphthalenes, tritolyl phosphate, organic esters, ethereal oils, cellulose plastic softeners, synthetic resins or varnishes may be added before, during or after chlorination. In an example, 10 g. masticated crepe in 200 cc. carbon tetrachloride is mixed with 1 g. sulphur and heated with chlorine until the gel formed redissolves to form a mobile liquid and the product is precipitated by adding 100 cc. alcohol and washed in boiling water to give a white mass containing 32% chlorine, soluble in acetone and benzol to yield a transparent, colorless film moldable at 130° C. If the chlorination is stopped before resolution, the gel which rises to the surface being removed, washed with solvent, treated with boiling water and dried, the product will be a semitransparent, hard, tough, elastic substance moldable at 130-140°.

De-Polymerization of Rubber German Patent 599,405

Rubber can be de-polymerized to give 40-60% solutions by treatment in suspension or solution with 10% of its weight of 53% nitric acid. A paste is first prepared by stirring 10 kg. rubber in 90 kg. benzol, whereupon 1 kg. of the 53% nitric acid is stirred in and the de-polymerization interrupted at the desired stage by neutralization with ½ kg. barium carbonate.

The de-polymerized rubber solution is decanted off and concentrated if necessary by evaporation. Coatings of this form of rubber are somewhat tacky but this defect can be remedied by a partial re-polymerization (immediately after the neutralization stage) with antimony trichloride or phthalic acid in alcoholic solution.

Cork-Rubber Composition British Patent 425,699

Rubber	100	lb.
Cork	100	lb.
Sulphur	3	lb.
Zinc Oxide	5	lb.
Stearic Acid	2	lb.
Mercaptobenzothiazole	0.5	lb.
Zinc Isopropylxanthate Pipe	eri-	
dine-l-Carbothionolate	0.5	
Paraffin	5	lb.
Nonox S	1	lb.
Lithopone	25	lb.
Chromium Oxide, Green	15	lb.

Cork Composition Canadian Patent 348,152

A mixture of phenol 13, paraformaldehyde 8 and diethylene glycol 30 parts by weight is heated to 210° F., 6.4 parts by weight of a 16% solution of caustic soda is added as a catalyst, and the heating is continued at about 210° F. until a sample of the liquid taken off will set in 10 minutes in boiling water. The product is immediately mixed with ground cork in the proportion of 80 lb. of the liquid and 150 lb. of cork particles. The treated cork is placed in a mold at about 300° F., where the reaction is completed and the comminuted cork is agglomerated into a cohesive mass of the desired shape.

Coating for Rubber Goods British Patent 427,228

,
100 lb.
1-5 lb.
100 lb.
50 lb.
10 lb.
5-20 lb.
5 lb.
8 lb.
10 lb.
to suit

Water sufficient to give a final concentration of total solids of 45-50%.

Thermoplastic Hornlike Rubber German Patent 615,050

Treat rubber with 70% hydrofluoric acid for 24 to 48 hours.

Rubber Curing Solvents Formula No. 1

Carbon Bisulphide	50 gal.
Petroleum Naphtha (140-220° F.)	50 gal.
Sulphuryl Chloride	1 gal.
No. 2	-
Carbon Tetrachloride	50 gal.
Petroleum Naphtha (140-220° F.)	50 gal.
Sulphuryl Chloride	1 gal.

Fire-Resistant Rubber U. S. Patent 1,966,271 Formula No. 1

A solution of 100 parts ammonium chloride; 6 parts ethylene glycol, and 3 parts glue in 300 parts of water is added to 3 parts of an antioxidant comprising a mixture of the condensation products acetaldehyde with a and \$\text{A}\$-naphthyl-amines, the antioxidant being wetted with a little alcohol. Sponge rubber is soaked in this solution and the excess squeezed out until the "wet" gain in weight is 120% on the dry weight. It is dried in a current of warm air. Sponge rubber treated in this way will not burn unless held continually in a flame, and one withdrawal from the heating flame, the sponge at once ceases to burn.

No. 2

Five parts casein are dissolved in ammonia solution, the bulk made up to 300 parts and 100 parts ammonium chloride dissolved in it. The sponge rubber is soaked in this solution and the excess squeezed out so as to leave in the sponge a quantity of solution equivalent to 120% of the weight of the dry sponge rubber. This is then dried in a current of warm air. The degree of fire-resistance can be adjusted by alterations in the proportion of ammonium chloride present.

No. 3

Excess selenium is boiled for 30 minutes with 20% ammonium sulphite solution and the solution obtained is filtered through glass wool. The sponge rubber is soaked in this solution and squeezed out until the increase in weight is 55% of the dry weight, and dried in a current of warm air. The selenium is slowly deposited spontaneously by exposure.

No. 4

Sponge rubber impregnated as in the preceding case with a solution of selemium in ammonium sulphite is exposed to an atmosphere of sulphur dioxide for liberation of the selemium. Alternatively finely powdered selemium is rubbed on to the surface and into the surface pores of sponge rubber so that some is permanently retained. The extent of fire-resistance depends upon the quantity of selemium retained.

No. 5

Sponge rubber soaked in a 20% solution of ammonium silico-fluoride is equeezed out until the gain in weight is 150% of the dry weight. It is dried in a current of warm air. Sponge rubber treated in this way will not burn unless heated continually in a flame, and on withdrawal from the heating flame at once ceases to burn.

No. 6

Thirty parts of finely powdered ammonium chloride are stirred into 100

parts of a 1% solution of rubber in benzene, and the suspension obtained is painted on to the surface of the sponge rubber. The solvent is allowed to evaporate and the surface is dusted with French chalk. The fire-resistance of the sponge rubber is markedly improved. A similar suspension of ammonium silicofluoride in a benzene solution of rubber has a like effect.

It is to be understood that the quantity of fire-preventing agent remaining in the pores is not sufficient to fill the latter in any case; that the porous structure is not changed and that the agent is deposited as a superficial coating on the inner surfaces of the pores.

Fireproofing Rubber British Patent 432,551

Five to fifteen per cent of any of the following is incorporated in the rubber: Triphenyl Phosphate Tricresyl Phosphate Triphenyl Borate

Rubber Calender Liner

The handling of miles of calendered sheet involves either efficient dusting methods, to permit rolling up of the sheet without risk of adhesion, or alternatively a good non-adhesive cloth that can be rolled up with the rubber. Where the sheet has subsequently to be cut into shapes and built up, its tackiness is important, so that dusting becomes out of the question in industries such as tire and footwar manufacture.

Gelatin	75 lb.
Glycerin, Commercial	85 lb.
Talc	30 lb.
Dye, Color to Suit	10 lb.
Water	800 lb.

The cotton is treated with this mixture on both surfaces and dried. It is then hardened by passing through a bath of 10% formaldehyde solution, dried, and pressed on a calender.

One thousand square meters of cotton sneet can be covered with 37.5 kg. geletin, 42.5 kg. glycerin, 15 kg. tale, 0.5 kg. dye, and 25 kg. formaldehyde.

Rubber Mold Lubricant

Sodium Hyposulphite	280	g.
Sugar	70	ğ.
Magnesium Sulphate Crystals	30	ğ.
Glycerin	15	g.

Hexamethylene Tetramine
Phenol
Sodium salt of the sulphuric
acid derivative of the reaction product of normal
butyl alcohol and a mixture of approximately 85%
ortho hydroxy diphenyl and
substantially 15% para
hydroxy diphenyl

5 g.

hydroxy diphenyl 5 g.

The composition thus prepared is added to substantially 20-30 times its weight of water. When applied on the surface of molds and press plates, which contact with rubber or other material to be vulcanized or molded, the film produced is markedly tough and resists rubbing off when the rubber or other material is pressed into the mold.

Lubricant for Vulcanizing Molds
Sodium Hyposulphite 3 lb.
Ammonium Carbonate 1 lb.
Water 97 lb.

Non-Adhesive Mold Liner

To a mixture of casein 45, glycerol 45, and kaolin 10 parts, add water to the required consistency. Apply 2-3 times on both sides of the cotton material. Dry 1-1½ hours. Then treat with formaldehyde. The total time is 6-7 hours.

Aqueous Latex Dispersions for Artificial Leather

A mixture composed of "smoked sheets" (rubber) 100, gasoline 200, olecacid 8, 25% ammonium hydroxide solution 20, casein 20, sulphur 8, zinc oxide 10, "Kaptax" 2, thiuram 1 part and water in accordance with requirements, produces stable emulsions when diluted with up to 50 volumes of water. A leather substitute of good physical and mechanical properties is obtained from rubber 100, rosin 19, oleic acid 5, wheat flour 15, glue 5, kaolin 10 and sulphur 5 parts.

Rubber Films and Threads

A hydrochloric additive product (1) is prepared, in 100% yield, by treating a 2% benzol solution of rubber with hydrochloric acid at 16.5-19° C.; after 15 hours the product is separated by precipition with alcohol. Glossy, transparent films may be prepared by spreading chloroform solution of (1) on a glass plate and allowing it to evaporate at 45-50° C. The films adhere to metals and may be

dyed; they may be combined with plasticizers, which reduce the strength but increase the extensibility. Threads may be prepared by dry spinning from a 7% chloroform solution of (1). The material is not readily combustible, and is but little acted on by hydrochloric acid (concentrated and 2N), potassium hydroxide (20% and 2N), sosp solutions, or 4N-sulphuric acid. Decomposition is effected by treatment with concentrated sulphuric or nitric acid or by prolonged heating at 55-60°.

Hard Rubber Coating U. S. Patent 2,023,582

A method of applying a hard rubber coating to articles comprises mixing aubtratially 500 g, of smoked sheet rubber, 180 g, of sulphur, 2½ g, of diphenyl-guanidine, and 2½ g, of mercaptobenzo-thizole, dissolving the mixture in substantially 2500 g, of benzine, applying the solution to an article to form a coating and vulcanizing the coating.

Wire Insulation Compound

The following formula provides an insulating compound capable of extremely rapid vulcanization and yet one which, when mixed and applied in accordance with the process defined, does not vulcanize during the extruding operation.

anne danne me candana	1,0.00		
Smoked Sheet Rubber	22	g.	
Reclaimed Rubber (Boot		-	
and Shoe)	10	g.	
Reclaimed Rubber (Whole			
Tire)	10	g.	
Mineral Rubber	5	g.	
Whiting	44.7	g.	
Zinc Oxide	2.5		
Antioxidant	1.5	g.	
Sulphur	1	g.	
Softener (Pine Tar Oil)	3	g. #	1
Ultra-Accelerator	0.3	g.	,
This stock is adented for	conti	1110118	

This stock is adapted for continuous vulcanizing carried on at a high rate of speed. For example in coating No. 17 Brown and Sharpe gage drop wire with a coating 3/4*in, thick satisfactory results are obtained when the speed of travel is from 400 to 500 ft. per minute when using a vulcanizing chamber 100 ft. long. The corresponding vulcanizing periods for these speeds would be 12 to 15 seconds.

Molded Brake-Lining U. S. Patent 1,963,511

A lining which is non-absorptive of oil or water comprises asbestos 29.7, carbon black 7.7, barium sulphate 12.8, lead oxide 2.0, rubber 33.2, sulphur 4.6, and an aqueous suspension of a phenolic or other infusible resin 10 volume per cent.

Electrical Insulation for Cables

Satisfactory insulation is achieved by coating the cable with a vulcanized mixture of synthetic rubber 15, filler (kaolin, chalk) 40, and asphalt 45%.

Artificial Leather (Gralek)

A fabric is coated with

Gasoline	871/4	lb.
Zinc Oxide	8	lb.
Sulphur	3	lb.
Thiuram	11/2	lb.

and then with a mixture of

na men with a mi	xture or	
Rubber	100	lb.
Leather Dust	150–2 00	lb.
Zinc Oxide	8	lb.
Sulphur	2	lb.
Lampblack	20	lb.
Accelerator	11/2	lb.
Pine Tar	5	lb.
Gasoline	1000-1200	lb.

The final coating consists of dry casein pigment, formalin 6% of dry pigment and alizarin oil 10%. Finally vulcanize and varnish.

Transmission Belt Dressing U. S. Patent 2,001,582

Neatsfoot C Rubber	il	,	1 13	lb. lb.
			10	•

Printing Blanket Patented

Formula No. 1

\$ A flexible and resilient printer blanket having a smooth surface which is resistant to oils and repellent to inks is made by applying a chlorinated rubber coating over the ordinary printer blanket. The coating varnish comprises chlorinated rubber (20 to 40), benzene (10 to 75), a plasticizer (3 to 10). Another varnish may contain chlorinated rubber (30), xylene (25), tricresyl phosphate (5); while in the other example there are combined chlorinated rubber (30), benzene (30), dibutyl phthalate (6). When it is desired to use pigments or dyes in the varnish it is preferred that the pigment, such as carbon black, is first mixed with the plasticizer and then incorporated in the chlorinated rubber solution.

No. 2 British Patent 423,556

In a printers' blanket of the type comprising a fibrous base and an outer coating of, or containing rubber, which is surfaced or ground in the usual manner, the outer coating is obtained directly from an aqueous dispersion of, or containing, rubber and is vulcanized to the The fibrous base may comprise a plurality of superposed layers of fabric material, e.g., felt, that are bonded together by a rubber cement or latex adhesive, containing vulcanizing ingredients, so that on subsequent vulcanization of the rubber coating the adhesive is also vulcanized. A preferred latex composition comprises rubber (as latex of 65.7% solids content) 100, formalin 4.65, water 79.75, potassium hydroxide 0.90, antimony sulphide 20, sulphur 6.8, whiting 75, ferric oxide 12, zinc oxide 2, sodium isopropyl naphthalene sulphonate 0.975, glue 0.375, heptaldehyde aniline condensate 1.5, acetone diphenylamine condensate 0.75 and solvent naphtha 1.5 parts; the coatings are dried at 90° C. and vulcanized at 135° C.

Mending Rubber Goods

Apply to the surface of the object a thin solution of rubber in benzol such as is used for sticking patches to auto tubes and allow a few minutes to evaporate solvent. Apply a generous coating of latex rubber and allow to stand a few hours. Can be used for mending auto tops, cuts in tires, hot water bottles, etc.

Rubber Packing Rings for Grooved Cans
In grooved containers with rubber
packing rings the caps are set in place
and the rings heated to 150-180° F. under
pressure for about 1 sec. The formulation of the rubber ring is of importance
for the proper speed of melting and the
proper degree of hardness. A typical
formula is (in percentages by wt.): rubber 14.10, balata 4.70, heavy spar 55.56
and chalk 25.64.

Puncture Proofing Tire Tube

A self-healing inner tube structurally designed to prevent deflation after puncturing is secured by lining the tube during its manufacture with a tread ply of rubber of special softening composition. The following formula gives satisfactory results:

Phosphoric Acid 2 lb. Clay 1% lb.

3 1 lb. Rosin Oil 93¼ lb. Rubber

The particular softening agent used is ortho-phosphoric acid of 85% strength. The clay serves as a vehicle for the phosphoric acid. The clay and acid are mixed together before being added to the other ingredients. The rosin oil serves as a softener and tack producer. The ingredients are mixed on a rubber mill in the usual manner and may be calendered and slit into strips. In the construction of an inner tube by the pole or flat drum method one of these strips is used as a lining for that half of the tube toward the tread. The application of heat to the tube results in vulcanization of the body structure, but the special stock layer, due to the presence of the chemical agent and absence of sulphur, accelerator, or other vulcanizing ingredients in its composition, does not vulcanize. On the contrary it becomes extremely plastic, almost viscous in form, and interiorly is very sticky. Although the non-tacky layer in the tube causes the surface of the special stock layer to be somewhat less sticky so that it will not adhere to the opposite wall of the tube should it come in contact therewith, it is preferable that the finished tube be kept in lightly inflated condition. In the event of puncturing by a nail the sticky layer adheres to the nail so that when the nail is withdrawn, it draws back some of the sticky stock with it so as completely to seal the hole through the body structure.

Puncture Proofing Tires German Patent 589,394

Use is made of mixtures of latex with animal, vegetable or mineral oils. A typical mixture contains ammoniacal latex 40, sesame oil 50 and olein 10%. The mixture is introduced through the air valve of the tire, distributes itself over the inner surface and automatically seals any punctures which may develop.

Gas Generating Composition for Rubber Balls

A stable mixture of ingredients from which to prepare pellets for use in in-flating hollow balls, etc., follows:

40 lb. Ammonium Chloride Sodium Nitrite 59 lb. Anhydrous Sodium Carbonate 1 lb.

The main constituents, viz., the ammonium chloride and the sodium nitrite, are commercial materials not completely dried. When maintained at 60° C., this gas producing mixture decomposes roughly 25 to 30 times more slowly than pellets prepared from dried materials but without sodium carbonate, and over 100 times more slowly than pellets prepared from undried commercial materials, again without sodium carbonate. They undergo no appreciable decomposition at ordinary temperatures, or is their value duminished for inflating rubber balls at 100° C. (212° F.) or over.

Rubber Vulcanization Accelerator U. S. Patent 1,963,084

Turpentino 100 oz. Sulphur 15 oz. Heat at 120-130° C. for 12 hours.

Forms for Molding "Bakelite"

Graphite Powder Clay Magnesia Cement Asbestos Powder Magnesium Chloride Mold to a paste.

Dental Thermoplastic Molding Composition

U. S. Patent 2,020,311

Twenty-five parts of rosin are melted or fluxed with 1 to 5 parts of glycerol (depending upon the abietic acid content of the rosin), preferably under a reflux condenser, and from 10 to 25 parts of aluminum stearate added to the mixture while it is still at a relatively high temperature, that is, 250° C. or there-above. From about 5 to 10 parts of rosin oil are then added, if desired, and after this has been thoroughly incorporated into the body, it is allowed to cod to a temperature of about 150° C., whereupon from 1 to 5 parts of triethanolamine stearate is added. Thereafter, wood flour may be incorporated. Prior to the addition of triethanolamine stearate, the composition, although elastic, is extremely sticky and gummy and unsuited for dental purposes.

Dental Impression Jelly

14 g. Agar-Agar Water Glycerin 10 minims 12 g. Kaolin

Dissolve agar-agar in water by heating in a pressure cooker for 11/2 hours. Then stir in other ingredients.

Plastic Molding Composition U. S. Patent 1,969,146

Phenol Formaldehyde Resin Charcoal, Powdered	4 lb. 6 lb.
Wood Flour	3 lb.
Pine Tar	1/2 lb.

Capsule Composition	(Cneap)
Potassium Silicate	
(30-33° Bé.)	70 g
Water Soluble Dye	2 2
Water	28 g

Capsule Composition

Gelatin				27	
Water				42.	7 g.
Allow to	swell	over	night	and	war
antly with	******	n~	4:1		

gently with stirring until uniform.

Glycerin (28° B6.)	10 g.
Water Soluble Dye	2 g.
Water	18 g.
Preservative	

Manufacture of Casein

"Rennet Casein" suitable for making Galalith and similar plastics is best obtained as follows: To fresh skim milk at 35° C. add sufficient rennet to effect coagulation in 15-20 minutes; stir 5-10 minutes and warm to 65° C. at the rate of 1° per minute; decant twice with water at 25°; drain and press out as much water as possible, disintegrate the press cake and dry at 43-45° C.

Plastic Composition French Patent 781,749

A composition for making pipes contains asbestos 85, fluid resin 15, lithopone 0.15, muldrite 1500 and cellulose 2200 kg. or vegetable fibers 85, resin 10 and rubber or latex or bitumen 5 kg.

Plastic Display Composition

Compositions based upon pigmented linseed oil, castor oil, and a non-alkaline thickening agent such as corn starch, have recently been suggested as constructional naterial for display work. They are also eminently pitable for coating theat-rical drop curtains and the like. They can be produced in various colors, and of a consistency permitting easy stencilling. For a yellow compound, 16 oz. of a

paste pigment in the ratio of 6 lb. white

lead to 4 lb. chrome yellow are worked up into 80 oz. of spar varnish, 10 oz. boiled linseed oil, 10 oz. Japan drier, and 2 oz. castor oil. Sufficient corn starch is then incorporated to yield a mass with the consistency of thick mortar, which is allowed to mature in the open air for about 12 hours before packing into airtight containers. Castor oil is an essential ingredient, since it assists maintenance of the solids in suspension for a very long period if the containers are air-tight.

When making up a bright red or orange composition in which the pigments accelerate drying, the above formula must be modified to the extent of using more castor oil (3 to 5 oz.), more lin-seed oil and less spar varnish. On the other hand, the slow-drying black compositions will require a higher proportion of varnish and japan, and as little as 1/2 to 1 oz. castor oil. This type of composition appears to be suitable for producing numerous figures required in industrial display work, the advantages being maintenance of flexibility and toughness after drying, good adhesion to supports and resistance to chipping.

Modeling Clay Formula No. 1

What is called molding compound by some artists is made by mixing two parts by weight of kaolin or powdered soapstone, which must be bone dry, and one part by weight of wheat flour, stirred into three parts of melted white beeswax (not too hot), and well kneaded before the wax cools. The mass may be colored to suit. A good modeling clay can be made from dry clay, mixed with glycerin instead of water. The mixture must be thoroughly mixed.

No. 2	
Plastic Clay	46 oz.
Cup Grease	24 oz.
Paraffin Wax	11 oz.
Rosin Oil	1 oz.

Polishing Plastics

Cast resins polish to a high, permanent luster. Rough cuts are usually ground, using the same type of equipment as required by wood or brass. Sand paper, garnet paper, belts or fine abrasive wheels are used. For most work, a generous supply of water is recommended, when wheels are used, to prevent over-heating and to keep the wheel clean. Surfaces which show tool or finding marks are given a smooth surface, preparatory to final polishing, by "ashing," in which an ordinary buffing wheel, made of muslin discs, of 12 to 14 in. diameter, is used. Wet pumice, kept in a shallow pan under the wheel so that the buff just touches it, is used as a polishing agent. Often, additional wet pumice, taken from the trough, is applied by hand or trowel above the piece being worked. Polishing is usually done, on larger pieces, by a second wheel, using bar wax or specially prepared polishing compounds. These wheels, usually 12 in. in diameter, operate around 1800 r.p.m. A third, clean dry wheel is used to give a final polish.

Tumble Polishing

For large quantities of small and medium sized pieces, tumbling is often used. Here, barrels of hard wood, lined with leather or heavy felt and operating at about 50 r.p.m. are used. Solutions vary with the article being polished, a common procedure calling for preliminary tumbling in dry pumice, to which wooden shoe pegs or similar agents have been added to provide friction. The pumice is later washed off and a second tumbling follows in damp hard wood sawdust. Other materials are sometimes used as well as pumice. A final operation consists in tumbling with powdered stearie acid or red oil. In some cases emulsions of carnaubs wax are used.

PROPERTIES OF NATURAL RESINS

	n					Direct
	Per			Boften-		Acid
	cent	Direct	Indirect	ing	Melting	Number
	Mois-	Acid	Acid	Point	Point	After
Natural Resins	ture	Number	Number	°C,	°C.	Running
Genuine Bold Pontianak	1.5	123	133	108	141	95
DBB Soluble Copal Chips	2.4	139	157	90	119	97
No. 1 Brown Kauri	5.4	57	67	120	152	3 5
Bold Black Scraped	1.5	20	36	125	164	17
Batu Bold Scraped		18	33	132	180	15
Pale Bold E. I. Singapore	0.7	20	37	128	156	9
Hard Dark Amber Congo	0.7	102	123	104	200	78
Congo Gum, Ivory Rescraped		92	111	91	144	92
Medium Pale Congo	0.4	110	132	85	220	70
Boea Medium Dark	2.9	126	149	115	148	95

Softening Point determined by the capillary tube method.

Melting Point determined by the Mercury Method Ranguswami, reported in the Journal of the Oil and Color Chemists Asso., 1930, Vol. 13, Page 287.

CLASSIFICATION OF NATURAL RESINS

- I. Low Acid Number Resins, including Damar and East India type.
 - A. Damar Resins-oil soluble-indirect acid number 25-45 M.P. 90-110° C.
 - 1. Batavia
 - 2. Sumatra
 - 3. Pontianak
 - 4. Padang
 - 5. Singapore
 - B. East India Fossil or Semi-fossil Resins—oil soluble—indirect acid number 25-40 M.P. 125-180° C.
 - 1. Batu
 - 2. Hiroe
 - 3. Rasak
 - 4. Macassar East India
 - 5. Bold Black Scraped
 - 6. East Indian Singapore
- II. Resins of High Acid Number originating in the East Indies:
 - A. Pontianak—Fossil resins—oil and spirit soluble—indirect acid number 103— 140 M.P. 135-145° C.

B. Manila resins

- Soft or Menlengket resins—spirit soluble—indirect acid number 135– 160 M.P. 110-135° C. Macassar
- Half hard or Loba resins—spirit soluble—indirect acid number 140– 150 M.P. 115–120° C. Loba and Macassar Loba
- Hard fossil resins—oil and spirit soluble—indirect acid number 105– 120 M.P. 140-155° C. Boea-Loewoe-Pontianak
- III. African Fossil or Semi-fossil oil soluble—indirect acid number 110-135 M.P. 140-220° C.:

A. Congo

- IV. New Zealand fossil or semi-fossil resins—oil and spirit soluble—indirect acid number 55-70 M.P. 120-160° C.:
 - A. Kauri
 - B. Bush Kauri

Melting Points of Synt	hetic Resins
Amberol BS1	99-110° C.
Bakelite BR352	93-104° C.
Bakelite BR2072	80- 91° C.
Beckacite 1101	102-112° C.
Beckacite 1102	102-112° C.
Beckacite 1113	102-112° C.
Akco Resin, Hard	125-130° C.
Amberol F7	118-125° C.
Amberol 226	117-133° C.
Amberol 801	117-133° C.
Beckacite 1112	110-125° C.
Lewisol No. 1	120–125° C.
Paranol, Hard	115° C.
Paranol, Extra Hard	125° C.
Akco Resin, Extra Hard	
Amberol K-12-A	148-175° C.
Bakelite XR2963	138-150° C.
Beckacite 1100	127-142° C.
Beckacite 1106	127-142° C.
Lewisol N2	130-135° C.
Robert Rauh N2	135-145° C.
Q. D. No. 1	135-145° C.
Q. D. K.	140-150° C.

Hardening Rosin

Five hundred kilograms of rosin are melted in a kettle. Thirty-eight to 40 kg, of hydrate of lime are added at a temperature of 205° C, and the mixture is heated to 260° C. which causes the lime to dissolve and the mixture to clear up. The acid number of the hardened rosin amounts to half that of the colophony. In Germany the rosin is heated for some time at 175° C. Six per cent of calcium hydrate (produced from marblestone) with a magnesium oxide content of not more than 3% is then added. It is advisable to grind the calcium hydrate to a paste with a little lineed oil. The English process, which is usually

The English process, which is usually carried out in enamelled kettles consists of stirring 6% of calcium hydrate (marblestone material) into the rosin heated to 60-80° C., and it is claimed to be

possible by energetic stirring and careful operation to raise the lime additions to as high as 10%. According to another American process, 100 kg. of colo-phony are heated to 232° C. Six per cent of calcium hydrate is then gradually stirred into the melt within about 15 minutes and the mixture heated to 268° C. within another 15 minutes. The opinions regarding the most efficient process are thus very different. It is important to determine the most suitable percentages of lime hydrate to be added, since working by "feel" may easily cause the production of turbid material. A rosin of an acid number 145 requires the addition of 9.8% of hydrate of lime or 10.5% of zinc white. A rosin of an acid number 180 requires 11.9% of hydrate of lime or 13% of zinc white. However, the rosin must always be heated to 175° C. before adding the lime. Hydrate of lime as well as zinc white must be absolutely dry. The lime hydrate should be freshly slaked, free from carbonic acid and finely dispersed, and it is always advisable to grind this material with a little linseed oil. Most rosins require only 6% of hydrate of lime or of zinc white (Green seal) free of carbonate. It is also possible to add both materials at the same time, as for instance, 2% of zinc white and 4% of hydrate of lime. The zinc white is added at a temperature of 220 to 240° C., the mixture boiled clear, the hydrate of lime added and the mixture heated for some time at 275° C. If the hydrate of lime contains more than 3% of magnesium oxide, the melt thickens.

According to the Haines Process, the lime rosin can be boiled directly with oil, satisfactory results being obtained with two different methods of application: The oil is either boiled with the whole of the roein at once or only with part of it, the remainder being added later

in form of a lime rosin-benzene solution 1:1. The results of this process are as follows: The viscosity of the pigmented varnishes decreases if larger quantities of the rosin are boiled directly with the oil. Skinning of the pigmented varnishes decreases in the same manner. The larger the quantities of rosin directly boiled with oil the more pronounced is the whitening of the varnishes in touch with boiling water and the slower is the disappearance of the whitening. The behavior of the products towards cold water is similar. Maximum adhesion after 24 hours of storing in water is exhibited by a varnish, half of the lime rosin contents of which had been boiled with oil. This varnish also exhibits the largest pressure resistance; it is also superior in its behavior towards rapid weathering while if subjected to normal weathering conditions, the gloss of the clear varnish decreases directly with the increase of the amount of lime rosin boiled directly with oil.

The larger the quantity of lime rosin boiled with oil, the more pronounced be-comes the sensibility of the product towards subsequent covering of the film with nitrocellulose lacquers.

The Koehler process for the direct boiling of lime rosin with oil is carried out as follows: The necessary quantities of rosin are disintegrated and dissolved at 80 to 100° C. in benzene (crystal oil). At a temperature of 105 to 110° C., 4 to 5% of hydrate of lime, free of carbonic acid and lumps, and suspended in ben-zene, is added. The kettle, or boiler, must not be filled to more than onefourth of its capacity since the process is accompanied by strong foaming of the contents. The temperatures must not rise above 120 to 125° at the most. After foaming has subsided, the varnish can be produced at once by adding acid-poor, water-clear stand oil (with an acid number of not more than 20), tapping the If white mixture and centrifuging. enamels are produced, lime rosins must be employed which are made from excelsior rosin. One day after the production of the varnish 17 kg. of zinc white and 18 kg. of lithopone are added per 60 kg. of varnish. The mixture is thoroughly stirred and left to stand for at least 2 days. Five kilograms of varnish and 2-3% of cobalt siccative are then added and the product thinned in accordance with requirements. The subsequent addition of varnish tends to improve the gloss of the product. One to 2% of gloss-improving substances may also be added if necessary. If top grade enamel

varnishes are to be produced it is advisable not to add linseed oil stand oil alone, but also about 20% of wood oilstand oil. However, both types of stand oil are to be boiled separately since if the two oils are boiled in common, the wood oil would thicken before the linseed oil had been boiled sufficiently.

Investigations towards improving the hardening process have led to the following formulae: 100 kg. of colophony are heated with 1 kg. of cadmium oxide to 200-250° C., stirring continuously. After complete solution, 5 kg. of hydrate of lime are added and the product left to cool down to room temperature. very satisfactory lime rosin varnish is obtained by this process.

Another process is the following: 0.5 cc. of 33% caustic soda solution and ce. of 33% causic som sommon and 45 g. of paraformaldehyde are added to 100 g. of crude cresol heated to 80 to 100° C. As soon as the paraformaldehyde has been dissolved, the mixture is cooled and added to 800 g. of colophony heated to 200-250° C. The mixture is then stirred until the smell of phenol has disappeared. One hundred grams of this alcohol-soluble product is treated with 1 g. of precipitated or fused lithium resinate, the product obtained being easily soluble and free of separations.

A number of important guiding rules have to be observed in the production of glycerin-rosin esters. Esterification is almost universally effected in apparatus with reflux coolers, the operating temperature being about 250° C. The amount of glycerin added exceeds by about 3% that determined by calculation from the acid number of the resin. Esterification is complete after about 3 to 5 hours. The temperature is then increased to 800-320° C, in order to drive off the excess glycerin, the water of reaction and the volatile constituents of the resin. It is recommended to add 0.5% of boric acid which accelerates the esterification and prevents re-saponification by the water of reaction.

Investigations carried through in the State Industrial Research Laboratory at Tokyo (Japan) resulted in the following discoveries: (1) If aluminum kettles are employed, this metal appears to exert a catalytic influence on the process of esterification. (2) The acid number of the resulting rosin esters drops rapidly if operations are carried on at a temperature of 200° C. (3) Fifteen to 19% of the rosin is the most suitable glycerin contents. Higher glycerin contents tends to soften the product. (4) Excessively long heating causes darken-

ing of the product. (5) Dehydrating agents increase the speed of esterification. Suitable dehydrating agents are the hydrates, oxides and carbonates as well as the organic salts of metals, for instance, the formates of calcium and barium. Undesirable additions are boric acid and manganese borate. (6) A metallic salt addition raises the rosin ester

softening temperature.

Typical and characteristic variation of the esterification process can also be observed in the various countries. In America, glycerin-rosin esters with acid numbers up to 3 are produced in aluminum kettles. "WW-rosin" is used for light colored products. After charging the kettle it is hermetically closed and the contents melted either in a vacuum or by passing through carbonic acid. Ten to 18% of glycerin (calculated on the amount of resin used) is then added and the mixture heated for some time to 205° C, and finally to 288° C. The water vapors are permitted to escape through a reflux cooler, the glycerin flowing back into the kettle. If rosin ester of an acid number of 5 to 10 is employed, about 12% of glycerin is added. It has also been found here that an excess of gly-cerin tends to soften the product while excessively long heating darkens the product. The varnishes produced from glycerin ester exhibit a high gloss. They are neutral in character and resistant toward basic pigments. They do not tend to crystallize, they are free of water, flow well, but are not easily mixed with drying substances, while colophony absorb them with ease. Balm and wood colophony, as well as mixtures of the two types of colophony can be esterified. The products of wood colophony are somewhat cheaper and exhibit a lower melting point. Instead of glycerin, other hydroxyl compounds, such as naphthol or benzyl alcohol, may be used, while fossil resins can be employed instead of colophony.

If, during esterizing, up to 10% of previously melted Congo or Manila copals are added, the melting points are considerably raised and the color darkened.

In Russia, rosin esters of an acid num-ber of 4 to 5 are produced by means of catalysts, such as zinc. Rosin and zinc catalyst are jointly heated to 275 to 280° C. Eighteen per cent of glycerin is then added, the product having an acid number of 4. If catalysts are not added, it is possible by adding 24% of glycerin to obtain a product with an acid number of 25.5. A. Kogan recommends zinc chloride as zinc catalyst, while another suitable catalyst is iron trichloride in connection with hydrochloric acid gas. original saponification number of about 173.3 is lowered by the catalytic process to about 30-40. The rosin is not appre-

ciably changed by the use of catalysts.

According to U. S. Patent 1,771,044, it is possible even to produce rosin esters of an acid number 1 by esterizing the rosin or the resinous acid with dichlorhydrine or dibromhydrine in presence of alkalies. For instance, 75 parts of WWcolophony are dissolved in 100 parts of alcohol containing 10 parts of caustic soda. This solution is heated to 80° C. (reflux cooling) and gradually treated with 25 parts of dichlorhydrine of a boiling point of 174° C. The mixture is then boiled 15 hours (reflux cooling), the sodium chloride produced is separated and the dichlorhydrine excess distilled off. The yield consists of 70 parts of rosin ester having a melting point of 74° C. and the acid number 1.

An interesting French process provides for the use of wax alcohols. Natural colophony brands or resinates, hardened colophony or synthetic resins are made to react with wax alcohols, such as cetyl alcohol or cholesterol. For instance, 85 parts of colophony, 15 parts of lanolin and 2 parts of hydrate of lime are processed together. After heating the mixture of the first two constituents to 200° C., the hydrate of lime is added in small portions, and under continuous stirring, and this temperature maintained for some time. The product of reaction is transparent; it is soluble in the common solvents and yields varnish films of considerable plasticity and resistance. Another variation of this process provides for the heating of a mixture of 88 parts of colophony and 12 parts of cetyl alcohol or cholesterol to 200° C. with, or without, catalysts. Conditions are improved by operating under pressure or in an inert gaseous atmosphere. Or 85 parts of colophony are heated with 8 to 10 parts of glycerin and 5 to 7 parts of purified lanolin.

Esterization can be combined with the rosin production in the case of ester rosins as well as in operating with lime rosins. For instance, colophony is melted at 193° C., 24% of wood oil added and the temperature raised to 250° C. per cent of glycerin is then added and the temperatures maintained at 288° C. for 6 hours. The kettle is finally removed from the fire and the glycerin rests removed by the addition of 5% of boric acid which forms volatile compounds with the glycerin. The product, thinned with lacquer benzene, represents a satisfactory varnish.

Glycerin rosin ester-wood oil varnishes must be boiled and cooled down rapidly. Cooling can be effected by means of cold water or by adding cold varnish or cold linseed oil refined with alkali. Boiling with 2.5% of litharge requires the consideration of the following factors: The varnish is water resistant and impervious to gases only if boiled at 296-302° C.; the fatter the wood oil varnish, the more durable is the film, but the more pronounced is the danger of gelatinization during boiling, which can, however, be reduced by adding colophony; the lower the degree of acidity of the ester rosin or the wood oil, the greater is the danger of gelatinization and the more difficult is the addition of drying substances; the larger the ester rosin contents, the brighter and harder is the film and the more rapid is the rate of drying. Attention is called to the fact that the addition of colophony has a slightly deteriorating influence on the quality of the varnish. Addition of linseed oil, fish oil, soya bean oil, etc., lowers the water resistance of the film and reduces the speed of initial drying, but improves the gloss, the life and the elasticity of the film. Ester rosin varnishes must never be mixed with cold oil varnishes, as the components of this mixture do not combine with each other in the cold.

Increasing the Melting Point of Rosin

The melting point of rosin can be raised from 61 to 91° C. by 2 hours blowing with air in the presence of 1½6 cobalt exide when molten, and to 107° C. after 6 hours. The amount of petroleum etherisoluble substances (hydroxy acids) increased from 17.89 to 46.07%, the acid number and saponification number decreased from 159.56 to 145.26, and 170.00 to 161.29 respectively, and the esterification number increased from 10.44 to 160.3.

Purification of Rosin

The rosin is crushed, melted in a kettle, allowed to stand for 30-60 minutes, decanted from the impurities into a second kettle, boiled 1 hour with 20% of a 9° Bé. sodium chloride solution; the supernatant sodium chloride is siphoned off, and the treatment with sodium chloride repeated till a sufficiently light colored rosin is obtained. Soaps made from such

rosin are lighter in color than those made from unpurified rosin.

Synthetic Resin Emulsion U. S. Patent 1,999,715

One hundred parts, by weight, of ground resorcinol are placed in a metal container or kettle which is jacketed with an outside container, in turn equipped for steam heating and water cooling. permits of temperature regulation during the chemical reaction as it is very essential to carefully control the temperature of the reacting substances in order to prevent their conversion to the insoluble stage. The kettle may be equipped with suitable agitators to permit the rapid churning of the kettle contents. To the resorcinol, 112 parts, by weight, of 37.5% formaldehyde solution (formalin) are added, and the temperature is increased to a maximum of 60° C., so that the resorcinol will dissolve.

In a separate container, 8 parts, by weight, of para-nitraniline are dissolved in 20 parts, by weight, of crosel having a boiling point range of from about 215 to 230° C. The melted para-nitraniline is added to the formuldehyde solution, and the mass is thoroughly agitated, the reaction temperature being raised to from 70 to about 75° C. A plasticizer of vegetable or animal oils and wax with a filler and suitable coloring material may be added as a paste to the mixture in the kettle. As an example, 8 parts, by weight, of clay, 0.8 part, by weight, of iron oxide, all best ground in a paint mill or ball mill to obtain thorough dispersion. The paste has been found to mix readily with the thickening liquid in the kettle.

For best results, the temperature should be maintained at no higher than 80° C. When the mass in the kettle becomes stringy, and before gelatinization can take place, the emulsification enhancing agent is added. A water solution (65 parts, by weight) containing 0.1% of bornx is added, first slowly, and then rapidly to the agitated resin. The borax. or any other suitable alkaline salt, serves to so enhance the emulsifying action of the wax and/or the oils (and gums, if any are used) as to enable them to properly maintain the resin in suspension in the water. This results from the fact that the borax or other alkaline salt used reduces the surface tension of the water from its normal value at the operating temperatures, thereby more readily effecting a wetting of the resin particles

and enabling the wax or other fatty material used to more easily retain the resin particles in suspension. The temperature is dropped to about 20° C. and thickening of the resin takes place. A solution of alcohol (about 65 parts, by weight) or an equal quantity of a 20% benzol or toluene solution in alcohol may be added to the emulsifier to obtain proper consistency. The varnish is immediately strained to remove any foreign particles. The mass is then cooled, with accompanying increase in viscosity of the varnish, and additional alcohol or solvent mixture may be added to obtain the desired viscosity.

Flexible Synthetic Resin U. S. Patent 1,999,097

Diethylene Glycol 106 oz. Phthalic Anhydride 148 oz.

These ingredients are mixed together and heated gently in a suitable receptacle until all of the phthalic anhydride has melted, the temperature of the mix is then gradually raised to approximately 165° C, and maintained at this temperature for approximately 4 hours. The resulting resin on cooling is a viseous liquid having a light amber color and is soluble in acetone, alcohol, chloroform, nitrocellulose and cellulose acetate solution.

Synthetic Molding Resin U. S. Patent 2,010,225

A mixture of 9 lb. of asbestos, 3 lb. of shellac and 2 oz. sulphanilic acid is repeatedly passed and repassed between hot rolls maintained at a temperature sufficient to keep the shellae in the composition molten. When the mixture in this manner has been rendered uniform in stribution, the resulting plastic mass may be pressed into slabs, or so-called biscuits of any desired type or shape. These biscuits may then, if desired, be subjected to a heat curing or baking process. The exact details and conditions of any such intermediate step will, to a large extent, depend upon the nature and service to which the later manufactured article is to be put. This product may then be placed in a mold either in biscuit (by softening on a steam table) or in a powdered form, and subjected to required heat and pressure necessary for forming a hard, resistant, less fusible object. Where a limited amount of agent and previous heat treatment of the biscuit material has been employed, it may be necessary to cool the mold during the pressing operation. The molding may take place in a number of ways but good results may be obtained by softening the biscuit material at 300° F., placing a slight excess in the mold and subjecting the same to 2700 lb. per sq. in. The pressure is not released until the material has cooled to a temperature sufficiently low to be readily handled without deformation.

Synthetic Resin Paper Size Emulsion U. S. Patent 2,022,004

Resin A

Glycerol 15.6 oz. Phthalic Anhydride 20.18 oz. Stearic Acid 64.22 oz.

The ingredients are heated together with stirring in a suitable vessel, the temperature being carried to 200° C. over a period of 1 hour, then maintained at this point until an acid number of 47 has been reached. This requires approximately 2½ hours.

The preferred method of converting the resin into an aqueous emulsion, defined here as a dispersion of very fine particles of the resin in water, is as follows: 100 parts of Resin A at 100° C. and 61.0 parts of a 5% solution of sodium hydroxide at 60° C. are added simultaneously and in proportionate rates to 349 parts of water at 60° C., with rapid agitation during the mixing operation. The alkali solution should be added slightly in advance of the resin, and the emulsion should be stirred for a few minutes after the mixing operation has been completed. This gives a 20% emulsion of the resin. The amount of sodium hydroxide used in preparing the emulsion is insufficient to neutralize completely the titratable acid in the resin. The resin is therefore not present in the water in complete solution, but as an emulsion, i.e., it is largely in the form of a physical dispersion in the water. This is a very substantial difference from those cases in which the resin is completely neutralized, as in the prior art. The suitability of the present emulsions enables one to use resins which are carried to a lower acid number and, hence, a more complete resinification. Lower acid numbers and higher resinification are necessary to give the improved water resistance when applied for the purposes of this invention. High acid number resins require alum in addition to alkali to develop their maximum water resistance; the use of more completely esterified products obviates this disadvantage. The emulsion can be diluted with warm water to any desired concentration.

Plaster of Paris Synthetic Resin Casts British Patent 425,742

Plaster of Paris casts are impregnated with an aqueous solution of the reacting components of phenol formaldehyde resins in the early or molecular stages of condensation to increase their hardness, toughness and gloss, and to secure their impermeability to water. The product is capable of taking a high polish, and of being stained. Instead of phenol, cresol and homologues thereof may be used. In an example a plaster cast is immersed until saturated in a mixture of equal weights of commercial cresol and 40% formaldehyde and 1 or 2 parts of a 50% solution of potassium hydroxide. The solution is warmed to 35° C. The object is then stoved at 100° C.

Synthetic Dielectric Resin Canadian Patent 342,586

Abietic acid 800, glycerol 770, phthalic anhydride 852, ethylene glycol 905 and linseed oil acids 80 parts by weight are heated under reflux to 175-180° C. for approximately 30 minutes, and 320 parts by weight of tung oil is added in 4 parts. Succinic acid (1820 parts) is added and the mass cooked until a resin is formed. The excess of glycerol is removed by vacuum distillation. The resin is used in coating compositions for fibrous material as cloth and paper in order to impart a flexible, tough film of good dielectric value, unaffected by mineral oil or petroleum or aromatic solvents.

"Albertol" Type Synthetic Resin

Formaldehyde 0.85 l.
Phenol 1 kg.
Hydrochloric Acid 0.02 kg.
Reflux 2 to 3 hours.

Pour off liquid and dry residue in vacuo at 100° C,

To 0.3 kg. of above resin add 0.7 kg. rosin and heat to 120-130° C. When solution is complete add 0.4% calcium oxide and heat to 290° C. Maintain at this temperature until a sample is soluble in oil and has an acid number of about 30.

"Haveg" or "Prodorite" Type Materials

An acid proof material suitable for tanks and other apparatus is made of

80% sand, an appropriate amount of coal or oil bitumen and of 5% acid resistant minerals (grog, clay, etc.); the mixture is leasted to 150-200° F. and molded to the desired shape. It sticks to iron, is resistant to hydrochloric acid and to diluted nitric acid. Commarone tar can be used as a protecting varnish for low temperature and for molded objects of a low mechanical strength. "Haveg" from asbestos and bauxite has a mechanical strength similar to that of cast iron.

Sound Record Composition British Patent 408,969

A particularly suitable resin is formed by the conjoint polymerization of vinyl chloride 80 and vinyl acetute 20%. The resin may be mixed with a filler, e.g., wood filler, cotton flock, silica, mica with a plasticizer, e.g., dibutyl phthalate, tricresyl derivative, glycol, glycerol esters.

Gramophone Record Composition

Composition
15 oz.
1.5 oz.
19 og.
19 oz.
5.5 oz.
40 oz.

For cheapness, part of the carbon black is often replaced by mineral black. The scrap is spew and rejected records,

the scrap is spew and rejected records, etc. The amount of lac varies, dependent upon the grade used, it being generally considered that T.N. Orange is about the lowest that can conveniently be employed at present.

Vinyl Resin

Canadian Patent 352,766

Polymerizo following at about 40° C.:
Vinyl Chloride 80 oz.
Vinyl Acetate 20 oz.
Hexane 100 oz.
Benzoyl Peroxide 0.5 oz.

Vinyl Acetate Resin German Patent 615,995

Water	200 g.
Vinyl Acetate	200 g.
Hydrogen Peroxide (30%)	1 cc.
Soda Ash	1 g.
Heat at boiling point for 1	to 2 hours.

Bleaching Beewax

To)	
	Water Potassium Bichromate Sulphuric Acid (60° Bé.) Boil.	70-75 cc.
a.	Sulphyria Asid	15 g.
	(60° Bé.)	15-20 g.
(Boil.	

add

b. Beeswax, Molten

100 g.

Stir until color becomes greenish blue. Cool. Remove solution shortly before wax solidifies. Boil wax with clean water to remove acid.

Synthetic Beeswax U. S. Patent 1,983,672

Formula No. 1

Five hundred grams of a mixture of the higher paraffin hydrocarbons melting at 74-76° C. (Superla wax) is mixed with 10 g. of manganese cleate and oxidized in a glass reaction vessel at 130-140° C. by oxygen passed through the hydrocarbons by means of a tube with many small orifices submerged in the hydrocarbons. The oxygen is passed through the hydrocarbons at the rate of approximately 1/2 cu. ft. per hour. At the end of 144 hours the contents of the vessel has gained in weight about 20 g. It has an acid value of about 23 and an ester value of approximately 100. In physical properties this product closely resembles beeswax except it melts at a temperature approximately 10 degrees above the melting point of true beeswax.

No. 2

Two batches of 1500 g. each of the ozokerite wax ("Utahwax") with a melting point of 73° C. are mixed with Mut 1% of their weight of manganese cleate and then oxidized simultaneously in 2 flasks A and B. Dry oxygen at the rate of 30° cu. ft. per hour is passed into the flask A and brought into intimate contact with the hydrocarbon therein. The oxygen and the vapors coming off from the first flask A are passed through a soda-lime tower and then into flask B. The temperature of each flask is maintained at approximately 120° C., and after oxidation for 288 hours the reaction is discontinued. The product in each flask resembles commercial becswax and is suitable for use as a becswax substitute. The acid value of the product in flask A is about 25.8 and its ester value about 50.6. The product in flask as is about 25.8 and its ester value about 50.6. The product in flask

B has an acid value of about 46.7 and an ester value of about 56.6.

Raising Melting Point of Montan Wax U. S. Patent 1,966,168

Formula No. 1

Crude montan wax with a melting point of 80° C. is fused. Two-tenths per cent of calcium hydroxide suspension is added to fused wax while continuously stirring, the temperature being slowly raised up to 90°. Stirring is continued at this temperature for about half an hour. In this way the melting point of the montan wax is raised to 85°.

No. 2

Crude montan wax having a melting point of 80° is fused and 0.2% of calcium hydroxide is introduced at a temperature above 100° while continuously stirring until uniform distribution has taken place. After about half an hour treatment the melting point of the wax is raised to 85°.

No. 3

Crude montan wax solution obtained in the course of manufacture is mixed with 0.2% of calcium hydroxide, care being taken that uniform distribution takes place. After the hydroxide has acted for about half an hour the melting point of the wax raised about 5°.

"Hardened" Stearic Acid Wax Stearic Acid 75 oz. Magnesium Oxide 5.3 oz.

Heat with stirring for ½ hour at 130-150° C. Pour at lowest possible temperature.

Illumination Candles Paraffin (50-52° C.) 79 g. Stearin 19.5 g. Carnauba Wax, Bleached 1.5 g.

Wax Lighting Tapers
Paraffin Wax (40-42° C. or
42-44° C. 65-85 g.
Ceresin (58-60° C.) 30-10 g.
Beeswax 2-3 g.
Turpentine, Thickened 3-2 g.
Wick of loose cetter thready 30 t

Wick of loose cotton threads, 30 together for a size of 2-4 mm., wound on wire.

	Burning (Burgundy Pitch	% lb.
U. 8	3. Patent 1,	954,659	Rosin W.W.	1/2 lb.
Paraffin W	ax	49 lb.	Zinc Oxide	11/2 lb.
		ole Oil 51 lb.	Melt together the wax	
			and add the zinc oxide ald	wly with good
	Molded Can	ılla	mixing.	
			No. 2 Ozokerite	421/ Ib
U. E	3. Patent 1,	960,994	Beeswax	63 1/4 lb. 31 1/4 lb.
Beeswax		70 oz.	Graphite Powder	4% lb.
Stearic Ac		20 oz.	Grapinio Tonder	476 10.
Paraffin W		10 oz.	No. 3	
"Cellosolve	e''	1 oz.	Beeswax	85 lh.
			Burgundy Pitch	5 lb.
Seeling W	x for Cand	le Decorations	Turpentine	10 lb.
•	12 101 COM		No. 4	
Rosin		50 g.	Ozokerite	95 lb.
Ruby Shell	a.c	3 g. 1 g.	Graphite Powder	5 lb.
Gypsum		+ R.	N -	
			No. 5	
	Dental Wa	BX	Ozokerite, Green	33 lb.
Stearic Ac	id	1 lb.	Paraffin Wax	50 lb.
Paraffin Sc		2 lb.	Rosin W.W.	16 lb.
Glyceryl T		1 lb.	Petrolatum N. C	⅓ lb.
Carnauba '		2 lb.	No. 6	
Ethylene G	Hycol Glycei		Ozokerite, Brown	90 lb.
Stearate		2 lb.	Graphite Powder	2 lb.
			Pine Pitch	8 lb.
	Ceresin W	ax	Rosin Oil	1/4 lb.
Carasin wa		of a mixture of		-
ozokerite and			Insulating W	ax
Starting w	th pure vell	ow ozokerite and	Carnauba Wax	1 lb. 14 oz.
melting toge	ther in the	following pro-	Yellow Beeswax	4 oz.
portions with	paraffin w	ax gives the fol-	Venice Turpentino	6 oz.
lowing blend		•	Gum Obsidian	6 oz.
Pure Ozo-			Sulphur	2 lb. 8 oz.
kerite Wax	Paraffin		Cook until thoroughly t	
White	Wax		This wax should have a	melting point
М. Р.	M. P.	gives	of 285° F. and a flash po	int of 499° F.
75° C.	50° C.	Ceresin Wax	•	
4 oz.	1 oz.	M.P. 73.5° C.		-
4 oz.	2 oz.	M.P. 71.7° C.		
4 oz.	3 02.	M.P. 72.5° C.	Recording (Phonogra	iph) Wax
4 oz.	4 oz.	M.P. 69.7° C.	Formula No.	1
When pure	white ozok	erite is used the		
following res			Stearic Acid	84 lb.
Pure Ozo-			Melt and add slowly wi	
kerite Wax,	Paraffin		Litharge	81/2 lb.
White	Wax		Boil off water at 220-23	0° F. Stirring
M.P.	M.P.	gives	must be of such type to	prevent caking
75.7° C.	58.3° C.	Ceresin Wax	at bottom of kettle. Wl	ien solution is
4 02.	1 oz.	M.P. 74.4° C.	complete add slowly (by s	ifting in):
4 oz.	2 oz.	M.P. 73.2° C.	Soda Ash	7 lb.
4 02.	3 oz.	M.P. 72.5° C.	When a drop cools to a	
4 02	4 oz.	M.P. 72.0° C.	action is complete. Driv	re off all one.
			froth and water by heating	z up to 270° F.
731.		Wares	If a brown wax is desire	d add to above
Ele	ctrotypers'		Stearin Pitch	2 lb.
	Formula No			
Beeswax		5⅓ lb.	If a black wax is wante	
Paraffin W	ax	3 lb.	soluble nigrosine to brown	TOTHIUM.
			•	

010	IE CHEMICA	AL FORMULANT		
No. 2		Wax for (Wounded) Trees		
Distilled Montan Wax	60 lb.	Formula No. 1		
Litharge	41/2 lb.	Rosin	60 g.	
Soda Åsh	4 lb.	Alcohol	40 cc.	
Paraffin Wax	30 lb.			
Follow method exactly a	s in Formula	Melt up the rosin, add t	ne alcohol cau	
No. 1.		tiously. Stir until cold.		
		No. 2		
		Melt up:		
Shoemakers' Sewing	Wax	Rosin	15 g.	
Candelilla Wax	2 lb.	Linseed Oil	2 cc.	
Rosin	55 lb.	Turpentine (Thick)	1 cc.	
Burgundy Pitch	20 lb.	Yellow Beeswax	2 g.	
Rosin Oil	4 lb.	Melt together below 78°	C.	
Lard	3 lb.	Add:		
Mineral Oil (Heavy)	1 lb.	Alcohol	4-5 cc.	
			1 0 001	
Shoe Finishers' Black St	ick Wax	Fill into air-tight cans.		
Candelilla Wax	9 lb.	Non-Inflammable	T3/1	
Rosin	1 lb.			
Carnauba Wax (North		U. S. Patent 1,98	1,132	
Country)	32 lb.	Cellulose Acetate	100 lb.	
Oil Soluble Black Dye	6 lb.			
Carbon Black	1/4 lb.	Triphenyl Phosphate Diethyl Phthalate	10 lb.	
Paraffin Wax	1 lb.	·		
***************************************		Transparent Foil or F	ilm Rosa	
Black Padding Wa	ax	•		
Carnauba Wax (North		British Patent 41	1,411	
Country)	40 lb.	Cellulose Acetate		
Ozokerite (Green)	2 lb.	(Anhydrous)	100 lb.	
Paraffin Wax	58 lb.	Acetone (Anhydrous)	400 lb.	
Rosin	2 lb.	Diethyl Phthalate	16.7 lb.	
Oil-Soluble Black Dye	7 lb.	Diacetin	5 lb.	
		Triphenyl Phosphate	8.3 lb.	
Tree Grafting Wa	x			
Wool Fat, Neutral	22 g.	Polychromatic Printin	0	
Rosin	40 g.	U. S. Patent 1,999),549	
Ceresin (58-60° C.)	10 g.	Dextrin 10, glycerol 10,	goon 10 tale	
Beeswax	10 g.	10, naphthalene 0.5 and w	ater 16 parts	
Rosin Oil	18 g.	are mixed with a pigment.	acc. 10 pares	
	- B.	and and a pigments		

•]	RU	BBE	R, RES	INS	WA	XES, I	PL/	STIC	s				319
	Occurrence	Free in beeswax, montan wax, carnauba, also as cerotate in insect	Free in montan wax.	Free in beeswax and montan wax.		As tri-palmitin in palm oil and Japan wax; as cetyl palmitate in spermaceti; as myricyl palmitate	As laurin in coconut oil and Japan	wax. As myristin in coconut and palm- nut oils.	A.	Spermaceti.	As ceryl palmitate in opium wax,	As myricyl palmitate in beeswax, carnauba, sugar cane wax.	Carnauba wax.	Cochineal war.	In wool fat and sperm oil.	Plant cholesterol.
83	Soluble in	Warm Alcohol	Methyl Alcohol	I	Alcohol Ether	Alcohol Ether	1	I	Alcohol Ether Benzol	İ	Alcohol	Ether Alcohol	Ether	l	Ether	Dell'or
TYPE ALCOHOL	Specific Gravity at 15° C.	.836 at 79° C.	i	i	.847	.846	ı	1	.810	i	1	l	i	i	1 1	ī
HIGHER WAX	Melting Point	77.8° C.	83 ° C.	91 ° C.	70.5° C.	62.2° C.	43.5° C.	53.8° C.	50 ° C.	59 ° C.	79 ° C.	.c. 88	103 ° C.	103 ° C.	147 ° C.	134 ° C.
WAX TYPE ACIDS AND HIGHER WAX TYPE ALCOHOLS	Formula	сн₃[сн₂]₂₄со.он	СН ₃ [СН ₂] ₂₆ СО.ОН	$C_{30}H_{61}COOH$	$\mathrm{CH_3[CH_2]_{16}COOH}$	$\mathrm{c_{16}H_{32}O_2}$	$C_{12}H_{24}O_{2}$	C ₁₄ H ₂₈ O ₂	c_{16} H_{33} OH	С18Н37ОН	C ₂₆ H ₅₃ OH C ₂₇ H ₅₅ OH	$C_{30}H_{62}(\mathrm{OH})_{2}$	C2+H4S(OH)2	$C_{30}H_{60}(OH)_2$	C27H440H	
WAZ	Waxy Material	Cerotic Acid	Montanic Acid	Melissic Acid	Stearic Acid	Palmitic Acid	Lauric Acid	Myristic Acid	Cetyl Alcohol	Octodecyl Alcohol	Ceryl Alcohol	Myricyl Alcohol	Anonymous Alcohol	Cholesterol or Cholesterol Al-	cohol [Iso-Cholestero] (Isomeric)	Phytosterol

d din grb PHYSICAL AND CHEMICAL PROPERTIES OF THE COMMON WAXES		- 2:0-4:0 .2668	0 1.3-2.0 20 47.8 3.	- 12 33.3 5.7	35 - 10-20 29.0 4.7	48.0 31	3 51.5 29.3	I		- 15-20 11-15 3-3.5 - 73-83 56-64 .03	(Dist.) (Dist.) (I	100 0 - 0 ° 0 0 - Ca 5%	100 0 - 0 0 0 - 0	.7-15 5.0-16.0 .6-2.6 6-20 90 11-35		40 86-91 14.3-15 1.5 5 30 4277	1	%##_co	51.5 3.0-4.0 .5-30 0.5-1.0 53.5 124-	- 25-43 3-4.6 12.2 - ery high
'ROPERTI	Setting Sa Point No	- 206-216	60-63 90-101	06-08	63-68 50-65	80-87 67-88	80-81 82-93	- 150-160	- 100-150	70-80 30-45 74-127		- 0-1.3	76 1.3	- 219-237	i	92-99 —	- 88-93	- 77-79	-122-134	- 101-104
EMICAL P	M.P. °C.	41	62-66	58	9 02-99	83-84 8	81-83.5 8	ı	62-70	73-84 7		35-75	59-76	52-59	82.0	12	1	6	41-59	37-41
L AND CHI	Ref. Index M.P. °C.	1	1.440-75°		1.456-75	1.472-43°	1	1	1	l		1.4331-1.4450	1.4415-1.4464	1.4518	ı	1.459	1.462-25°	1.456-25	1.4198	1.480
PHYSICA	Sp. Gr.	.993997	.960947	.980	.972	.992998	.932970	i	806.	ı		.870910	.913923	.976993	ı	.879	.876	.878	.932963	.945
	Bayberry (Myrtlewax)	Not a true wax	Beeswax	Cane Sugar Wax	Candelilla Wax	Carnauba Wax	Chinese Insect Wax	Cotton Seed Wax	Flax Seed Wax	Montan Wax	D	Farann wax Not a true wax870910 1.4331-1.4450 26-56	Ozokerite Geresin Wax913923 1.4415-1.4464 59-76	Not a true wax	Raphia Palm Wax	Sperm Whale Oil: Head Oil	Body Oil	Arctic Sperm Oil	Spermaceti932963	Wool Wax

				RUI	BBER, I	RES	INS	, WA	XE	S, PLA	STICS			821
,	11		Solubility	in Fusel Oil	Soluble	1 4	Politica		1 :	Soluble	1		Soluble	Boluble
	25-200 .1-0.2	.5 .01	solubility in	Carbon Tet- rachloride	Soluble	1 7	eigniog		1 :	Soluble	1	l	Soluble	Soluble
	8	96-99.5	u.		••									
	Variable 130-186	86.		Solubility in Turpentine	Soluble	1:	Solubie		1	Soluble	I	1	Soluble	Soluble
	11	1	.5	1 4	_		solu-	aple		_				
	10 55–180	.5-30	AXES	Petroleum Ether	Soluble in hot Insoluble and cold	1	Cold—insolu- ble	Hot—soluble	١	Soluble	1	ı	Soluble	Soluble
034	5-2.0 5-15	0.5	MON W	lity in ier	in hot d	,	-nlosu	Hot-soluble	1	in id hot	ı	1	Soluble in hot and cold	Soluble in hot and cold
400	- 147-180	200	F COM	Solubility in Ether	Soluble ir and cold	1	Cold—insolu- ble	Hot-s	1	Soluble in cold and hot	•	•	Soluble in and cold	
SH. 1	ADCLIERANIS OF MAARES 30-60 — 198 .5-2.0 ver 100 — 147-180 5-15	i	O ATA	Solubility in Chloroform	Cold—insolu- ble Hot—soluble	1	Cold—insolu.	Hot—soluble	1	Soluble in cold and hot	ı	1	Soluble in cold and hot	Soluble in cold and hot
	ADCLT 30-60 over 100	49-56	ITY I				Cold	Hot		Solut cold				
•		1.4380 49	SOLUBILITY DATA OF COMMON WAXES	Solubility in Acetone	Insoluble in cold Boluble in hot	i	Insoluble in	Not very sol- uble in hot	ı	Insoluble in hot and cold	1	١	Insoluble in cold and slightly solu-	Insoluble in cold and sol- uble in hot
	. 1.07–1.08	1		Solubility in Hot Acetic Anhydride	on ta,	Bulloo		acetytated	ŀ	Dissolves and solidifies on	اه	Dissolves and solidifies on	8m1000	ı
	Hardened (Hydrogenated) Oil — Roein	Stearin		Esolubility in Alcohol		ς Ω.		•	Insoluble		Montan Dist. Mon- tan Wax 70° C.	1	Insoluble	4.
	(Hydra			u2	•	:	:		Insect	۲. تا	į	: 8		oeti
	Hardened Rosin	Stearin .			Beeswax	Candellila	Carnauba		Chinese Insect	Japan Wax	Montan	Ozokerite	Paraffin	Spermaceti

SOAPS, CLEANERS

bon	15, 0	BEANDIES		
		,	T- 0	
Solvent Liquid Soaps		Soap	No. 9	F 1
Formula No. 1		Ammonia (0.880)	5 kg. 25 kg.
	500 kg.	Cyclohexanol	,	10 kg.
Hexalin 250-3	800 kg.	Water		60 kg.
	99 kg.		o. 10	00 ng.
Water 12 No. 2	08 kg.	Soap		10 kg.
		Ammonia (0.880)	5 kg.
	00 kg.	Tetralin		10 kg.
	50 kg.	Water		75 kg.
	08 kg. 92 kg.	Other liquid soap	s can be mad	de accord-
No. 3	л ь.	ing to the followin	g formulae:	
	00 kg.	Formula .1	No. 11 No. 12	2 No. 13
1:1 Hexalin-Methyl Hexalin 2		Coconut Oil	21	6 kg.
	70 kg.	Soya Bean Oil	8	12 kg.
Water # 1300-18		Potassium Hydroxid		
The ingredients are stirred tog	rether in	Solution (50%)	9.5 4.6	9.6 kg.
an indirectly steam-heated pot		Sugar	12 8	— kg.
clear solution is formed; this	is tested	Borax Glycerin	2 — — 6	kg. 12 kg.
for alkalinity.		Potassium Car-	_ 0	12 kg.
Hexalin or methyl hexalin		bonate	_ 2	— kg.
partially replaced by other sol- shown below:	vents as	Water	55.5 71.4	
No. 4		Oil of Lavender		0.1 kg.
Linseed Oil 184	kg.	Linalyl Acetate		0.1 kg.
Hexalin 275		The oil is first	run into a r	oan fitted
Potash Lye (50° Bé.) 73	3.5 kg.	with an open stean	a coil which	serves to
Water 387		both heat and agit	ate the pan	contents.
Carbon Tetrachloride 80	kg.	Heat the oil to ab	out 70° C. s	and grad-
No. 5	¥	ually add the car		
Coconut Oil 51		until the oil is com		
Linseed Oil 42		It will be found a before all the alkal		
Hexalin 130 Potash Lye (50° Bé.) 42		This is one method		
Water 615		which is likely to	occur partic	ularly in
Carbon Tetrachloride 120		the case of cotton-s	eed oil and t	o a lesser
Similarly, equal weights of be		extent when cocons	ut or palm l	kernel oil
high-boiling petroleum distillates		is used. When sap		
used in place of carbon tetrachle		add sugar, glycerin	, etc., and f	inally ad-
No. 6		just the water cor somewhat, then add		
Soap	35 kg.	required.	color and p	criume ii
	10 kg.	Where possible i	t is an adv	antage to
	55 kg.	use soft water, as	salts of he	rd water
No. 7	00.1	result in the forma	tion of corre	esponding
	28 kg. 10 kg.	insoluble metallic so		leposit or
	60 kg.	give a cloudiness in	solution.	
	2 kg.			
No. 8		Liquid Son	ap Shampoos	,
	30 kg.	Liquid soap shar		
Trichloroethylene	25 kg.	from olive oil pota	sh soap dis	solved in
		hot 80% alcohol in		
	33	22		

soluble, although the solution becomes slightly clouded on cooling. Dissolve the soap (1 part) in alcohol (4 parts) in a vessel which can be heated on a water bath and so constructed that alcohol is not logt by volatilization. When completely dissolved add coloring matter and perfume.

The formulae given are only a very few of the many that are available. Even using the same constituents of a given formula, the number of combinations could be varied in relation to fatty acid content, etc. Obviously the relative percentages of oil and alkali required for saponification would vary only between narrow limits.

Production of Liquid High-Content Potassium Soaps

German Patent 613,224

Formula No. 1

Colein 350 g.	
Coconut Oil Fatty Acid, (free from Stearic	
(free from Stearie Acid) Distilled 50 g.	
Alcohol 150 cc	
Water 210 cc	·
b. Potassium Acetate 50 g. Caustic Potash (48° Bé.) 190 cc	
Caustic Potash (48° Bé.) 190 cc	

Mix the two solutions. Soap contains 40% free fatty acid, is liquid down to 0° C, and gives no jelly on standing.

110. 2		
Futty Acid of Low-Boil		
ing Fraction of Spern	1	
(Whale) Oil	1000 p	ŗ.
Cocoanut Oil Fatty Acid	l	
(Low Titre) Distilled	220 g	ζ.
Adinic Acid	75 (7.
Alcohol		
Caustic Potash (48° Bé.)	630 c	c.
Water	700 c	c.
	(Whale) Oil Cocoanut Oil Fatty Acid (Low Titre) Distilled Adipic Acid Alcohol	Fatty Acid of Low-Boil- ing Fraction of Sperm (Whale) Oil 1000 g Coconnut Oil Fatty Acid (Low Titte) Distilled 220 g Adipic Acid 75 g Alcohol 450 c Caustic Potash (48° Bé.) 630 c Water 700 c

Mix a and b, and add c, with stirring. Clear, liquid soap with 40% free fatty acid.

Liquid Soap (15%)

23.qu.u =02.p (2= /c/		
Coconut Oil	12	kg.
Castor Oil	4	kg.
Potassium Hydroxide		
(50° Bé.)	8	kg.
Water	76	kg.
Potassium Chloride	0.5	kg.
Sanonify with warming, all	me to	stan

Saponify with warming; allow to stand for 1-2 weeks, separate clear liquid by siphon, filter sludge through a Seitz filter, put both together; optionally use alcohol or glycerin.

Liquid Olive Oil Soap

Two hundred and twenty-seven kilograms of potash are dissolved in the minimum quantity of water, and into the solution is stirred a mixture of 182 kg. olive oil, 362 kg. palm oil and as much cocount oil previously warmed to 49° C. Alcohol is next run in (170 l.) and the liquid hented to 82° C. (under reflux it is presumed). After saponification and cooling, 5.6 l. water are run into the alcoholte soap.

Liquid Coconut Oil Soap

Six kilograms potash are dissolved in 20 l. water and the solution run into 20 kg. eccount oil warmed to 49° C. After adding 25 l. of alcohol the mixture is kept at 82° C. to saponify, when it is left to cool for 24 hours. Eighty liters of water are then added, with a little sugar, potassum chloride or glycerin if necessary.

Glycerin Liquid Soap

Thirty-five parts of good soft soap are well mixed with 21 parts glycerin, and 7 parts of water well crutched in. This is followed by 14 parts alcohol. This solution is subjected to a fairly long sedimentation after adding tale or punice. If excessively alkaline it must be first corrected by the addition of oleic acid. Perfuming or coloring can be done if desired.

Liquid Scaps

Coconut Oil	10	kg.
Castor Oil	5	kg.
Lard Oil	2	kg.
Canstic Potash (31/2 parts		.,
solid)	161/2	kø.
Water		suit

This should be easy to make. Warm up the mixed oils and add the caustic solution. Heat gently. When clear and bright, like syrup, add sufficient distilled water to the consistency required, using phenolphthalein solution (½%) to correct.

Another mixing that will not lather as readily as the previous one, but which has the advantage of being an excellent cleanser, the power of which is only slightly diminished even in hard water, is as follows:

Lard Oil, Olein or Castor Oil 50 kg.
Glycerin 150 kg.
Caustic Potash Solution
(38° Bé.) 20 kg.

Carbonate of Potash Dissolved in 5 parts of Hot Water 3 kg.



This can be per following should	fumed	slightly and	the
following should pleasing, result:	give	a delicate,	yet

iensing, iceuit.	
Lavender Oil	2 kg.
Bergamot Oil	1 kg.
Geranium Oil	1 kg.
Patchouli Oil	1/4 kg.

About 1% of this should be sufficient to give the desired effect. The method of making the above soap should follow along the lines described and should present no difficulty.

Formaldehyde Soap Solution

Soft Soap	40	lb.
Alcohol	30	lb.
Formaldehyde	20	lb.
Distilled Water to make	100	lb.
As to perfume, oil of lavend	der (1	aboı

ut

Liquid Disinfecting Soap

a. Coconut Oil Soya Oil	18	kg.
Soya Oil	2	kg.
b. Caustic Potash (38° Bé.)	12	kg.
c. Water, Soft	68	kg.
Potassium Chlorida	0.5	le ce

Mix a, saponify with b, dissolve in c. Prepare:

Turkey Red Oil (70%) 3 kg. d. Phenyl-p-Hydroxy-Benzoate 20-25 dg.

The solution d is enough for 100 kg. of above made soap-base. Add perfume.

Disinfectant Scrub Soaps

Cheap disinfectant soaps in England ordinarily consist of suitable tar acid derivatives emulsified in a solution of rosin soap. Creosote, phenols, cresols and naphthalene are the usual disinfectant agents. The following directions are for the preparation of liquid disin-fectant soaps suitable for scrubbing floors, etc.:

Formula No. 1

Ground Rosin	17 lb.
Caustic Soda, 30%	3 lb.
Water	5 gal.
Crude Cresol	3 gal.

Boil the caustic soda in 1 gal. of water and add the rosin gradually to this. When dissolved and partly saponified, add 2 more gal. of water with continuous boiling and stirring. Add 2 gal. of cresol with stirring, then the remainder of the water and cresol. Keep covered until cold.

No. 2	
Water	61/2 lb.
Powdered Rosin	3%, lb.
Powdered Soda Ash	
Powdered Naphthalene	1 lb.
Filtered Creosote	% lb.
Soft Soap	⅓ lb.
D: 1	¼ lb.

Dissolve the soda ash in water and heat to boiling. Add the rosin and heat until saponified. Mix the soft soap and naph. thalene separately and add the creosote to this. Add the mixture to the rosin soap with continuous stirring.

Pine Oil Sambhian G

Time Oil Scrippin	g Boap
Corn Oil Soap	50 lb.
Pine Oil	10 lb.
Diglycol Laurate Alcohol	5 lb.
	3-5 lb.
Mix until uniform.	A transparent

jelly like product is formed.

Liquid Pine Oil Soap

Formule No. 1

Pine Oil	300 kg.
Soya Oil Fatty Acid	100 kg.
Water	60 kg.
Varmed gently to be liquefied,	

Clear Soap Oil, 1 part mixes with 40 kg. Turpentine 4 parts or

Benzoline 4 parts or

Carbon Tetrachloride 4 parts

Dichloro-Ethylene 4 parts

Naphtha 4 parts to clear oils, which give excellent emul-

sions in water (1:1 to 1:2). Above made Pine Oil Soap 12.5 kg.

Pine Oil 12.5 kg. Spindle Oil, Refined.

2° Engler, at 50° C. 75 kg. yields clear oil, gives excellent emulsions with water.

No. 2

Rosin WW_F/G 15 kg. Soya Oil Fatty Acid 30 kg.

Pine Oil 105 kg.

Take off 40 kg. and keep aside. To the remaining 110 kg. add: Water 135 kg. Caustic Potash (50° Bé.) 15 kg.

20	-	
	-	

ir until glassy-transparent	, add the
To the product add	
Water less than	a 300 kg.
(a tough, jelly like soap	paste)
or Pine Oil	100 kg.
(water soluble, liquid son	
Pine Oil Jelly Soa	n
Soya Oil Fatty Acid or	r
Linseed Oil Fatty Acid	40 kg.
Pine Oil	25 kg.
Warm gently.	0
Add:	
Water	15 kg.
('austic Potash (50° Bé.)	8 kg.
Caustic Soda (36° Bé.)	12 kg.
Water (optional)	15-30 kg.
No. 4	
Pine Oil, "Soluble"	
Soya Oil Fatty Acid or	
Linseed Oil Fatty Acid	25 kg.
Pine Oil	35 kg.
Warm gently in	v
Water	10 kg.
Caustic Potash (50° Bé.)	10 kg.
Pine Oil	160 kg.
Time On	20
Pine Oil Cleaning Pa	
Glycol Laurate	5 lb.
Pine Oil	25 lb.
Mix and add to following w	hile stirring
vigorously	EQ. 11.
Water	50 lb.
Caustic Soda	¼ lb.
Soap Paste Paint Cle	aner
Soap Chips	20 oz.
Mineral Spirits	10 oz.
Water	69.3 oz.
Oil of Sassafras	0.7 oz.
This is a semi-solid or	heavy soap
paste, white and permanent.	It is very
effective as a cleaner for t	oainted sur-
faces. It is also used as a	cleaner for
carnets and rugs. The SORT	i is anowed
to soak in the water which is to bring all the soap int Same is then agitated vigor	tnen neared
to bring all the soap in	to Boldrion.
the mineral enjoints is added	nd then the
the mineral spirits is added a oil of sassafras.	ing then the
OH OF SESSETIES	
Waterless Soap	
Oleic Acid	4 lb.
Turpentine Substitute	1 lb.
Tarbentine onostruce	0 11

Neutralized with a solution of caustic potash (1:1), 2 of water added to form

Industrial Spirit Castor Oil

2 lb.

1 lb.

a	paste	and	15%	οf	powdered	borax	in-
co	rporat	ted.					

Soap Powders				
Formula No. 1				
Palm Kernel Oil Fatty Acid 3 lb.				
or				
Tallow)				
Hard Fat Futty				
Hard Fat Futty Bone Fat Acid 2 lb. Palm Oil (Bleached)				
Palm Oil (Bleached)				
Caustic Soda (36° Bé.) 3 lb.				
Soda Ash 12 lb.				
No. 2				
Soft Soap Fatty Acids 6-7 lb.				
Hard Soap Fatty Acids (as				
above) 4-3 lb.				
Caustie Soda (37° Bé.) 6 lb.				
Water 50 lb.				
Soda Ash 36 lb.				
No. 3				
Soft Soap Fatty Acids 12-15 lb.				
Hard Soap Fatty Acids 8-5 lb.				
Caustic Soda (37° Bé.) 12 lb.				
Water Glass (36-38° Bé.) 6 lb.				
Soda Ash 30 lb.				
Water 32 lb.				
No. 4				
Soft Soap Fatty Acids 18 lb.				
Hard Soap Fatty Acids (as				
above) 7 lb.				
Caustic Soda (37° Bé.) 15 lb.				
Water Glass (36-38° Bé.) 8 lb.				
Soda Ash 25 lb.				
Water 27 lb.				
11 14 45 4				

Soap Flakes

To make high grade soap flakes, a To make high grade soap flakes, a good quality charge consisting of 75% tallow and 25% coconut oil, with or without the addition of 2% or less of rosin, should be used. The mixture should be loiled and finished as for toilet soap, then chipped and dried. Care must be taken in drying in order to produce. then chipped and dried. Care must be taken in drying in order to produce a uniform chip and avoid overdrying. The temperature of the soap chips should never fall below 30° C.; the temperature of the finished flakes should be between 40 and 45° C. The flakes should be milled twice to give transparency and polish. The most satisfactory shape to avoid breakage of very thin flakes is the square. square.

Soap for "Soap Noodles"	,	
Coconut or Palm Kernel Oil Tallow or Hard Fat	28 4	g. g.
Caustic Soda (38° Bé.)	10	

Potassium Carbonate	
(30° Bé.)	10 g.
Water	10 g.
Salt Solution (24° Bé.)	10 g.
Sugar Solution (24° Bé.)	10 g

Borax Soans

Soap from Kettle	1000 lb.
Powdered Borax	130 lb.
Lye (40% Caustic Soda)	23 lb.
Perfume, etc.	sufficient
The seen in sun into the	

The soap is run into the crutcher, the bornx, etc., added, and the whole crutched until the materials are thoroughly mixed. The physical condition of the soap is of less importance than when the soap has to cool in the frames and, therefore, the incorporation of larger quantities of bornx becomes feasible.

Various methods are available for the manufacture of soap powders, fillers being introduced before or after the soap is converted into powder. In the former case spoken of as the "continuous" process, the soda ash used takes up the

case spoken of as the "continuous" process, the soda ash used takes up the excess water present with the sonp, forming hydrated carbonate of soda and thus obviates the necessity of drying. A sonp powder of this type suitable for laundry and general purposes can be obtained from the following formula:

| Borax Soap Powder | Soap | 42 lb. | Soda Ash | 42 lb. | Powdered Borax | 15 lb. |

The soap is run hot from the kettle into the crutcher, and after thoroughly mixing with the soda sah and the borax, it is run over chilling rolls to chill the soap and crystallize the salts. The product is scraped off the rolls, the coarser particles being ground further. Alternatively, the mixture, after leaving the crutcher, is allowed to season for a few days, after which it is ready for powdering and packing.

Washing Powder

Fatty Acids	27.7-45.4 kg.
Sodium Perborate	4.8-13.5 kg.
Soda Ash	17.1-23.2 kg.
Water Glass	Ü
(Dry Basis)	0.6- 2.4 kg.
	•

Abrasive Washing Powder

Soap Sodium Sand	Carbonate	5 -10.2 5.6-10 73.7-81.5	kg.
			-0.

Washing Powder Formula No. 1

Cut into small pieces

Hard Soap Waste	10	kg.
Dissolve in Water	46	kg.
Add		Ü
Water Glass	10	kg.
Sodium Carbonate, Calcined	39	kg.
Mix well to obtain homogene	ous	mass.
No 2		

Hard Soap Waste Water	20 kg. 41 kg.
Water Glass	9 kg.
Sodium Carbonate, Calcined	35 kg.
No. 3	J
Hard Soap Waste	40 kg.

water		38	Kg.
Water Glass		4	kg.
Sodium Carbonate,	Calcined	35	
to get a 20% powder.			•

Note: the sodium carbonate is added only partially to the formulas 1, 2, 3, % is put on the bottom of the mixer before starting. Blow air into the warm mixture. Let cool for 24 hours.

Ammonia Washing Powder

Hard Soap	Powder (Alkaline)	1	lb.
Ammonium	Carbonate	1	lb.

Household Scourer

Colloidal Clay	1	lb.
Silica Floss	1	lb.
Alkaline Hard Soap Powder	4	lb.
Silicate or Carbonate of Soda	1	lb.

Fermentative Washing Powder
Sodium Carbonate 75 g.
Bile, Precipitated on
Kieselguhr 25 g.
100 g. of this powder are applied to
50 kg. laundry batch.

Cold Processed Soap British Patent 403,500

A method for preparing "cold processed soap" is to stir a mixture of 170 lb. of palm kernel oil with 9 gal. of 36° Bé. caustic soda solution. In a separate container, 6.5 gal. of a mixture containing equal parts of palm kernel oil and rosin is heated to 250° F., cooled to 110° F., and quickly added to the first mixture. After stirring for 10 seconds, the soap is run out through a valve in the bottom of the mixing pan, and subsequently treated in the usual manner.

Addition of rosin makes a more satisfactory and standard product than is usually obtained by cold process methods.

Cold-Process Carbolic Soap

For toilet purposes a cold or semiboiled soap is used, which retains the glycerin liberated from the fat. The following is a typical formula:

Formula No. 1	
Coconut Oil	80 lb.
Tallow	40 lb.
Soda Lye (38° Bé.)	60 lb.
Phenol	3 lb.

The fat and lye are thoroughly stirred at 35° C. until combination occurs and the soap is streaky. The phenol (dissolved in a little water) is crutched well into the soap; perfuming is sometimes done with a little clove, lavender or rose-mary oil. When cold the soap is cut into tablets and wrapped in air-tight package.

No. 2		
Bone Fat	150	lb.
Rosin	150	lb.
Carbolic Acid Solution	25	lb.
Caustic Soda Live (37° B6.)	150	lb.

The rosin and fat are melted together, and when the temperature is about 75° C. the carbolic acid is stirred in. The mixture is then added to the lye gradually, heating until the reaction is complete. The soap is framed and cooled and cut into bars of the usual size.

Cold Process Soap British Patent 432,227

Cold-process fat-resin soaps are made by treating fatty matter with just suffi-cient alkali for saponification, treating a mixture of rosin and fat or oil with alkali sufficient to saponify only the rosin, mixing the two products, and adding alkali to saponify the surplus fat. For example, 100 lb. of palm-kernel oil is stirred rapidly with 4.5 gal. of 36° B6. caustic soda for 10-15 minutes, 4 gal. of a melt of rosin in an equal weight of palm-kernel oil is treated at 110-135° F. with 0.5 gal. of 36° Bé. caustic soda, the products are mixed, and immediately 1 gal. of 36° Bé. caustic soda is added, and the mixture stirred for a few seconds and run quickly into the frames, where it sets and saponification is completed.

Dry Cleaner's Soap British Patent 407,088

Soaps for use with dry-cleaning solvents, especially carbon tetrachloride or

trichloroethylene, consist of a fatty acid soap with a content of a polyglycol, with or without a chlorinated aliphatic hydrocarbon. An illustration is the following: 14.2 g. of sodium hydroxide is dissolved in 25 cc. of water and stirred into 100 g. of oleic acid and 100 cc. of trichloroethylene. Next 70 g. of triethylene glycol or 50 cc. of diethylene glycol is added. The product is dissolved in trichloroethylene.

Soaps Containing Pine Oil German Patent 616,029

Formula No. 1	l
a. Pine Oil Caustic Potash	100 g.
	12.5 g.
b. Coconut Oil Fatty	

18-25 g. Acids Treat a at 80-100° C., neutralize the

product with b. No 2

Pine Oit	100	g.
a. Pine Oil Caustic Soda (95%)	4	ğ.
b. Fatty Acid	19	g.

As in No. 1. Solid, water-free soaps, high transparency.

Solid Pine Oil Soap U. S. Patent 2,007,974

Take one part water and two parts olive oil soap containing about 10% of water in the condition of flake or powder and when those are well blended stir in about one or two parts of pine oil. The vessel containing the mixture is placed in a kettle surrounded by glycerin and the temperature of the soap, water and oil is gradually raised to about 240° F. by heating the outer kettle. Steam is given off causing frothing of the soap with a great increase in volume of the mass. While some oils ordinarily begin to volatilize below this temperature, the soap raises the boiling point and permits them to be completely merged and held. When the heat, frothing and stirring have secured a uniform mixture, the mass is permitted to cool and solidify.

permitted to cool and solidity.
The solid sosp lathers well, but slowly
and yields at all dilutions a perfectly incorporated oil. It has the pleasant odor
of pine oil but has the firm feel of
anhydrous soap. The well fixed character of the oil is proved by the fact that the soap does not render white paper greasy after long contact with it.

Medicated Soaps

These types of soap can be made in two ways, either milled or by the cold process; as to their efficiency for the purpose for which they are intended, opinions differ, some claiming that they are of no value, others that certain complaints can only be cured by their use. Certainly much can be said for the latter statement, particularly when the complaint is in the nature of a skin disease such as eczema, and even without the addition of a specific body, toilet soaps which are superfatted with bodies such as landin or petroleum jelly naturally have a beneficial action on the skin.

No compound in skin soaps can compare with the well-known ichthyol variety. This compound can either be incorporated with flowers of sulphur and camphor or it may be used alone. Two mixings are given below contaming these bodies.

The first examples given are of the milled variety, which is certainly the best form of tablet both from appearance and as giving a perfect blend of the various bodies.

Ichthyol and Sulphur

Soup Chips	28	lb.
Ichthyol	41/2	oz.
Vuseline	2	oz.
Zinc Oxide	2	07.
Flowers of Sulphur	2	07.
Chlorophyll	11/2	θZ,
Medicated Perfume	4	04.

Tehthyol

Soap Chips	28	16.
Ichthyol *	7	oz.
Vuseline	2	oz.
Medicated Perfume	4	07.
Zine Oxide	2	07.
Chlorophyll	11/4	ez.

The antiseptic value of the tublets is enhanced by the use of the medicated perfume, which gives the type of odor used in a well-known line on the market, having a ready sale as a medicated toilet soap.

Medicated Perfume

Eucalyptus Oil	18 cc.
Terpineol	18 cc.
French Lavender Spike Oil	18 cc.
Red Thyme Oil	8 cc.
Clove Oil	8 cc.
Peru Bulsam	6 cc.
Camphor	3 g.

The soap and additions are milled in the ordinary way; it may be found necessary to mill more than the usual three times on account of the liquid nature of the additions. This may be obviated somewhat by using the soap chips a little drier than the usual 76-77% fatty acids-say about 78-79%.

The chlorophyll used is the oil-soluble type, dissolved in a little medicinal paraffin, or if this is not available the perfume may be warmed slightly and used as medium.

All other kinds of medicated milled soups can be made on the foregoing principle, leaving out the ichthyol, etc., and adding whatever is needed; the percentage used varies from 21/2 to 5, the lower figure being more general.

The other variety is the well-known cold process soap, a very fine preparation for the feet. This sonp, owing to the case with which it is made, is one for the small numifacturer with his limited phart. It contains permanganate of potush, and the directions for its use are: Wash the feet and allow the lather to remain in contact with the skin a munite or so before rinsing. The instructions for its manufacture are as follows: Melt the tallow and coconut oil together, and at 120° F pour in the canstic sods in a thin stream, stirring all the time; add the perfume and then the water, keeping the mass continuously on the move. When the soap is of the consistency of cream, which should be only about 3 to 4 minutes from the start, ponr into a wooden frame and just crutch the permangamite solution here and there in the mass; do not thoroughly mix it in. The appearance obtained is similar to marble graining. After standing 15 hours, covered and free from dranghts, the block of sonp is ready for entting, the size of tablets being usually

The mix	ing	for	the	above	soap	is:
Tallow					80	11
I'mannut	0.1					

Canstie	Soda,	669	Twaddell	80		lb.
Water				28		1b.
Perfum					1	lb.
Perman	ranate	of	Potnah in			

1000 cc. Water

Perfume

Pine Uil	1	cc.
Cassia Oil	1/4	cc.
Lavender Spike Oil		cc.
Patchouli Oil		cc.
Ditolyl Methane		cc.

Another soap made as above, leaving out the permanganate and using in its place stavesacre seed oil with a different perfume, is also sold for the removal of head vermin in children, and may be included in the list of medicated soaps,

Perfume

Sassafras Oil	5 cc.
Geranium Oil	1 cc.
Sandalwood Oil W.I.	2 ec.
Terpineol	5 cc.

The active principles of the last-named soap are the stavesacre seed oil and the sussafras oil—a very effective combination. These few examples embrace the whole range of medicated soaps, the only alteration in other cases being the medicating substance, the percentage of which, as mentioned before, ranges between 2½ and 5.

Antiseptic Scaps

An odorless phenolated soap can be made by mixing in about 3% of a fatty acid phenol ester such as phenyl stearate, palminate or olente. These esters are non-irritant to the skin and stable to alkalies. Iodine has been used in soaps. It does not have a very active antiseptic action when in the form of its compounds and is therefore employed as a solution in alcohol or in potassium iodide. Iodide is not stable however, as may be seen from the fact that soaps containing it change from brown to a light yellow in a short time. A better way of introducing rodine into soap is to add it in the form of a compound with an unsaturated acid such as oleic. A large number of so-called rodine soaps are made with potassium iodide and are quite stable, although they are not really iodine soaps.

Sulphur is a useful therapeutic for certain skin troubles. Its action is due to a mild antiseptic effect combined with reducing properties. Sublimed sulphin is generally used. The difficulty of getting sulphur into the water-soluble form may be overcome by using a combination of certain terpenes with alkaline sulphides and polysulphides. The solution of the clear brownish liquid in water gives a white emulsion with a slight alkaline reaction. It is non-irritant. A tar sulphur soap is widely sold for the treatment of a variety of skin diseases. It is a brown soap prepared by dissolving 2 lb. of potassium sulphide in a small amount of water, and adding 20 lb. of yellow stock soap together with 4 lb. of birch tar oil. The mass is milled several times.

The manufacture of soap incorporating mercury or corrosive sublimate is not an easy matter. The mercury salt reacts rapidly with the soap to form complex insoluble compounds. An improved process for incorporating mercury makes the soap contain an excess of

free fatty acid, which prevents the chloride from reacting with the sonp. In another process, the mercury salt is mixed with an alkaline casein solution, forming a mercury albuminate soluble in alkali. Mercuric iodide is used in some somes. It is best added by mixing 4 parts of mercuric iodide with 3 parts of potassium todide and 2 parts of water, then incorporating the precipitated salt with the milled soap. The method of using nonionized complex mercury compounds is one that shows promise. These compounds give no black precipitate on addition of ammonium sulphide in the cold. Those which give no precipitate on prolonged standing are the best suited for the purpose.

Germicidal and Antisept Coconut Oil Soap Base Cresol U.S.P. Mercuric Chloride 1-2000 Solution)	ie Sonp 50 g. 5 g. 45 g.
Iodine, Ichthyol, Camphe Formula No. 1 Sonp Base	ог Вопрв
Coconut Oil Ceylon Cunstic Soda (38° B6) Canstic Potash (38° B6) Lanolin Camphor No. 2 Iodine Soap	25 kg. 10 kg. 2 kg. 1 kg. 2 kg.
Same, but add Potassium Iodido n	1-1.5 kg.
Water, Hot No. 3 Ichthyol Sonp Same as No. 1, but add Ichthyol or Anmonium	2 kg.
Ichthyolsulphate Perfume	1-1.5 kg.
Peruvian Balsam Lavender Oil Cassia Oil Benzoin, Tincturo Perfume only for No. 2 or	120 g. 100 g. 100 g. 200 g. No. 3.

Boric Acid Soap Sapamin-Phosphute (100%) 10 oz. Boric Acid 5 oz. Glycerin 5 oz. Distilled Water 20 oz. Triethanolamine Laurylsulphonato 60 oz.

000	-					
Sand Soap				of rosin also assist		
Coconut or Palm Kernel Oil	20	kg.		of coconut oil is inci		
	11	kg.	воар	is required to lather	Treety	
Pumice, Finely Powdered	10	kg.				
Bolution of Benzoline,				Wool Scouring	Bath	
Tetralin	. 8	kg.	Oli	ve Oil Soap		40 lb.
Turpentine Oil in Turkey			An	amonia 28%		20 lb.
Red Oil (1:1)		ا بـ			-	
Perfume	0.5	%		Transparent Glyces	rin Sos	ns.
Mixture of				Formula No		
Lavender Spike Oil	5	cc.	_			
Rosemary Oil	4	cc.	a.	Prepare a solution		
Peppermint Oil	1	cc.		Caustic Soda (40°		20 g.
Caraway Seed Oil	1	cc.		Alcohol (90-92%)	14 g.
			l	Sugar		10 g.
Washing Tablets			Į	Water		11 g.
			i	Glycerin		11 g.
Formula No. 1			١.	Warm to 60-70° C.		
Perhorate of Soda	32	07.	b.	Add first melted		
Granulated Borax	35	oz.	l	Stearin, White		10 g.
No. 2			İ	then		•
Perborate of Soda	35	OZ.		Coconut Oil		10 ~
Borax	17.5		1	Tallow, White		18 g. 12 g.
No. 3	17.0	UZ.	1	Castor Oil		4 g.
	07		1	Castor Oil		* g.
Perborate of Soda	27	oz.	1	No. 2		
Borax	58	oz.	a.	Caustic Soda (35°]	3é.)	22 g.
No. 4				Alcohol	,	20 g.
Perborate of Soda	4	oz.		Glycerin		20 g.
Borax	12	oz.		Sugar		10 g.
No. 5			1	Water		10 g.
Perborate of Soda	34	oz.	l	Warm to 60-70° C.		•
Borax	18	oz.	Ъ.	Stearin		12 g.
Soda Ash	22	OZ.	1	Coconut Oil		20 g.
In each of above formulas n	aake	up to	1	Castor Oil		5 g.
100 with soap. Crutch with	son	; cut		37. 0		
into squares and dry.				No. 3		
			l	English Transpar	ent Sos	ap.
Wool Throwers Soar)		a.	Caustic Soda (38°	Bé.)	50 g.
Olive Oil Foots	12	lb.		Alcohol (90-95%)		50 g.
Corn Oil	46	lb.	l	Sugar		17.5 g.
House Grease	20	lb.	1 .	Water, 60° C.		23 g.
Soda Lye, 36° Bé.	3	lb.	b.	Pig Fat or Tallow		37.5 g.
Potassium Carbonate (Dry)	5%	lb.	1	Rosin, Pale		12.5 g.
Potassium Hydrate (Solid)	23	lb.	1	Coconut Oil		50 g.
			l			
Borax Laundry Son	3		F	'illed (Cheap) Trans	parent	Soaps
Finished Soap	1100	1 lb	1	Formula	No. 1	No. 2
Soda Ash		5 lb.	a.	Caustic Soda		
Solution of Carbonate of		, 10.	١ "	(38° Bé.)	77	48 g.
Soda (30%)	9,	5 lb.	1	Sugar	21	— g.
Solution of Metaborate of	-		1	Water	36	— cc.
Soda (s.g. 16)	2!	5 lb.	1	Filling Solution *	90	50 cc.
Silicate of Soda (40° Bé.)		5 lb.	1	Alcohol	12	20 g.
Soap Stock) lb.	b.	Coconut Oil	53.5	40 g.
The nature and proportions			1	Pig Fat or Tallow	53.5	40 g.
and oils are important. In			1	Castor Oil	42	20 g.
way the oils cottonseed, co	eonui	t. and	* 1	filling Solution,		•
palm-kernel, particularly the	las	t two	w	ater, boiled	800	cc. 200
mentioned, take up and hold fi	llers	better	Su	gar	51 52	g. 70 g. 60
than tallow and hardened oils.	Th	e pres-	Ba	tassium Carbonate lt	52	g. 40
		•				

Transparent Soap		Blenching Soda	
Hard Train Oil Fatty Acid	40 kg.	a. Water Glass, Commercial	
Sova Bean Oil Fatty Acid	60 kg.	(36-38° Bé.)	30 g.
Caustic Potash (50° Bé.)	42 kg.	b. Water	25 g.
Potassium Carbonate	13 kg.	c. Ammonium Carbonate	45 g.
Water	75 kg.	Dilute a with b, warm up in	a steam
		heated kettle with stirrer, add	e and mix
Filled Soap		to homogeneous distribution.	Pour hot
Palm Kernel Oil	900	on flat iron pans or on stone-f	loor, cool,
a. Tallow	200 g. 100 g.	turn with shovel, grind.	
Bone Fat	100 g.		
b. Water Glass	80 g.		
Tale	60 g.	Stain Removing Powde	r
o. Water	60 cc.	U. S. Patent 2,022,262	3
d. Caustic Soda (25° Bé.)	370 cc.	For removal of iron stains fr	om cotton
, ,		and rayon textiles.	om conton
Melt up a, keeping extra palm kernel oil. Add b molte	20 of the	Sodium Chlorite	1 oz.
tle to d, and boil to right of		Sodinm Oxalate	1 ez,
Add c as water-suspension.		Potassium Dihydrogen	2 (74)
salt water (23-24° Bé.) 8-1		Phosphate	2 oz.
test. If soap is too "sharp,	" add the	•	
remainder of the palm kerne	l oil until	_	
right. When tests show satis	factory re-	Dry Peroxide Bleaching P	owder
sults, boil 2 more hours at	id cool in	U. S. Patent 1,986,672	!
covered kettle.		A bleaching powder comprise	na nn nn-
		parently dry mixture obtainab	
		acting a hydrogen peroxide solu	
Soap Perfume		sodium bicarbonate and ther	
Cinnamic Alcohol	100 g.	anhydrous sodium carbonate a	
Neroli	50 g.	proportions of substantially 10	parts of
Petitgrain (Grasse)	50 g.	30 volume per cent of hydrogen	peroxide,
Orangeflower Absolute	10 g.	6 parts of sodium bicarbonate	
Hydrarom Fleur d'Orange	5 g.	parts of anhydrous sodium carb	onate.
Rose Otto (Bulgarian) Orris Concrete	15 g. 5 g.		
Costus (10%)	20 g.		
Sandalwood, E.I.	80 g.	Bleaching and Washing Po	wder
Bergamot	180 g.	French Patent 783,871	
Musk Ketone	40 g.	Formula No. 1	
Musk Ambrette	20 g.	Sodium Perberate	10 kg.
Coumarin	60 g.	Sodium Pyrophosphate	14 kg.
Vetiverol	70 g.	Soda Ash	8 kg.
Heliotropin	85 g.	Magnesium Silicate	1 kg.
Rhodinol, Pure	50 g.	**	
Methylionone, Pure	60 g.	No. 2	
Benzoin Resinoid	60 g.	Sodium Perborate	15 kg.
Phenylacetaldehyde (50%)	40 g.	Sodium Hexametaphosphate	10 kg.
****]	Soda Ash	9 kg.
Automobile Tar Solver		Magnesium Silicate	1 kg.
	,	Boap	49 kg.
Naphtha Ethylene Dichloride	40 oz. 90 oz.		
Diglycol Laurate	5 oz.	Stone, Brick and Masonry C	leaner
~- g. Jeor Laurate	J 044	TI C Det at 1000 000	-cont i

Automobile Cleaner

10 fl. oz. 2 pt. 1 pt. 6 pt. 1-2 lb.

Diglycol Laurate Kerosene Naphtha Water Kieselguhr

Stone, Brick and Masonry Cleaner U. S. Patent 1,990,383

Forty gallons of soap-bark extract formed from 9.5 lb. of soap-tree bark by steeping in water are mixed with rosin oil 1.25, raw linseed oil 1.25, an aqueous gum tragscanth solution (containing 1.25 oz. of the gum), (1¼ to 22%) hydrochloric seid 10 gal.

Brick and Masonry Cleaner Use a saturated water solution of ammonium bifluoride.

Deale Clauses

Diam Cicanei		
Caustic Soda, Powdered	15	oz.
Chalk, Powdered	25	oz.
Caustic Potash, Powdered	60	oz.
Keep dry and pack in air-ti	ght	tins.

Washing Compounds for Use in Canning The greatest surface is cleaned by a solution of a mixture of sodium hydroxide 2.8, soap 0.2, water glass 14.1 and sodium hypochlorate 4.8 (chlorine 2.3%)

but this has some corrosive action.

Cleanser for House Façades

Trisodii	ım Phosphate	75	g.
Sodium	Metaphosphato	20	g.
	Red Oil		g.
	Hydroxide		Ľ.
Water	to desired	concentrati	on.

Floor Bleaches

Oxalic acid has long been used to bleach or whiten discolored wood in its natural finish, especially floors. After applying this chemical, however, the wood is left so white that the spot usually must be stained lightly to restore it to the shade of the surrounding wood. Sodium perborate, which is sold in drug stores for use as a mouth rinse and a tooth powder, is a far milder blenching agent. Although one may have to rub the moistened powder on the discoloration a longer time than if an oxahe acid solution were used, the after effects are not so conspicuous. It is also particularly effective when mixed with equal parts of sodium metasilicate.

Cleanser for "Parquet" Floor Saponify

Caustic Soda (128–130°) Water	$\frac{6.64}{26.36}$	kg.
	4 5.15	

Alcohol, Denatured		45.4	ı.
The whole	poured into		

Trichloroethylene 900 kg. The product gives a stable emulsion with water.

Cleansing Preparation for Galoshes a. Carnauba Wax, Fat Gray 1 kg. Beeswax 0.5 kg.

ь.	Borax Capillary Syrup Water	0.5 kg. 0.3 kg.
M	Water elt up a, dissolve	25 1. b by short boiling,
dd dd	b to a and stir	until cooled, then
Th	inner (as above)	12 l.

0.5 kg

Olive Oil Soan

Trisodium Phosphate

Caustic Soda

Cleanser for Dishes, Glasses, etc. Formula No. 1

Sodium Metaphosphate	53 g.
Caustic Soda	2 g.
No. 2	
Trisodium Phosphate	55 g.
Sodium Metaphosphate	43 g.
Caustic Soda	2 g.
No. 3	
Trisodium Phosphate	75 g.
Sodium Metaphosphate	23 g.

Caustic Soda No. 4 Trisodium Phosphate (Monohydrate) Sodium Metasilicate (Peatahydrate) 40 g. Sodium Metaphosphate

Mechanical Dishwashing Preparation Sodium Metaphosphate 40 oz. Trisodum Phosphate 15 oz. 40 oz. Sodium Silicate Sodium Hydroxide 5 oz.

Glass Cleaners

Glass Cleaner in Cake Form Infusorial Earth, Finest

Powder 4 oz. Precipitated Chalk 2 oz. White Soap 2 oz. Boiling Water 2 oz.

Reduce the soap to fine shavings and dissolve in the boiling water. Then add powders which have been previously mixed and put through a fine sieve. Press into molds the size of the cake required and allow to dry.

White Soap	750 g.
Sodium Carbonate	20 oz.
Hot Water	120 cc.
Infusorial Earth	250 g.

Dissolve the soap (in fine shavings) in the hot water in which the sodium salt has been dissolved. Then add the infusorial earth in very fine powder. These soaps may be perfumed slightly by the addition of equal parts of oil of sassa-

	50m 6, C
fras and cedar oil to suit soaps get very hard in the time, owing to infusorial eart the property of absorbing co water.	
The following formula is example:	another
Powdered Pumice Stone Ammonium Oleate Ammonia (28%) to make Shake before using.	2 oz.
Ammonia (28%) to make	3 OZ.
Shake before using.	
Cleaning Mixture for Beer (
Use 1-3 g. per l. water of o mixtures (finely ground):	ne of the
Formula No. 1	
Trisodium Phosphate	600 g.
Sodium Carbonate	350 g.
Sodium Silicate	50 g.
No. 2	•
Sodium Carbonate	700 g.
Sodium Metaphosphate	300 g.
No. 3	
Trisodium Phosphate	800 g.
Sodium Bicarbonate No. 4	200 g.
Sodium Silicate	150 ~
Trisodium Phosphate	150 g. 850 g.
- The party	
Window Glass Cleaner	•
a. Mix	
Neuburger Chalk, Ppt.,	40
Finest Viennese Lime	20
Calcium Carbonate, Ppt.,	20
Heavy	25
Bolus, White	15
b. And grind with a mixture	of
Water	90%
Alcohol, Denatured	5%
Ammonia (sp. g. 0.91)	5%
Gun Cleaner and Solver	nt
	fl. oz.
	fl oz.
	fl. oz.
Butyl "Cellosolve" 1 Kerosene 4	l fl. oz. l fl. oz.
	oz.
Special Cleanser for Very Dir	ty Hands
Coconut or Palm Kernel Oil	
Fatty Acids Sova Bean, Linseed, Peanut	6 g.
Soya Bean, Linseed, Peanut Oil Fatty Acids	6 g.
Castor Oil Fatty Acid	3 g.
Pine Oil	6 g.
Alcohol	6 g.

Lanolin

Caustic Potash (50° Bé.) 6 cc.
Water 6 cc.
Pumice, Fine Powder until pasty
Citranella, ''Spike'' Oil,
Terpincol as Perfume to suit

Antiseptic Cleaner for Ice Cream

At the conclusion of the freezing operation drain the ice cream from the freezer. Ruise the strainer, hopper, and outside of the freezer particularly at the head, with cold water. Fill the freezer two thirds full of cold water, run one-half minute, and drain.

Fill the hopper full of water at 140° to 145° F, and add a half pound (1 cup full) of cleansing powder. Wash the strainer, hopper, and outside of the freezer with a brush. Drain the solution into the freezer (the freezer should be at least two-thirds full), run one-half minute, and drain the freezer.

Remove the head, scrub with a brush, being certain to clean out the front bearing. Wash the hearing ond of the dasher with a brush, remove from freezer and wash. Place dasher and head in sanitary place until used.

Before using the freezer, fill the hopper with water at 100° to 110° F., making certain that the screen is covered. Add sufficient chlorine to give 100 p.p.m. and stir well. If desired, the chlorine solution emilie pumped into the hopper from a special tank. Pour some of the chlorine solution into the front bearing, Place dasher in freezer and fasten the head in place. Drain the chlorine solution into the freezer, operate the freezer is then in excellent sanitary condition, except that the reur bearing may be contaminated, and is ready for use.

Lavatory Cleaner

One method is to add niter cake (acid acdium sulphate) to the water in the lowl. Another consusts of a mixture of sodium cuibonate (16 purts) and caustic soda (3 parts), and there are others depending on the liberation of chlorine.

A cleaner can be made up of sodium sulplate (88 parts), sulphuric acid (9 parts), and diatomaccous earth or some other fine abrasive material (3 parts).

Another suggestion is to mix powdered soap with four times its weight of powdered potassium carbonate.

Coconut Oil	10 lb.
Potassium Hydroxide	1 lb.
Sodium Hydroxide	1 lb.
Water	10 lb .

Dissolve the potassium hydroxide and sodium hydroxide in the water and mix with the coconut oil. Set aside in a warm place for a few hours to saponify. Test for neutrality and dissolve the product in 6 oz. of water. The resulting liquid soap does not cake and lathers freely when used in small quantities.

Laundry Bleach

Chlorii Washi Water	ng		0	11/2	lb. lb. gal.
		_			-

Allow to stand for a few days and filter.

Laundry Blue Good Quality

Formula No. 1

Ultramarine	60 lb
Bicarbonate of Soda	40 lb
Glucose	12 lb
No. 2	

Cheap Quality

Ultramarine	18 lb.
Kiln-Dried Blue Earth	20 lb.
Terra Alba	15 lb.
Bicarbonate of Soda	45 lb.
Glucose	10 lb.
No. 3	
Lime	5 oz.

Lime

Water 10 oz. Stir until smooth and mix with a hot solution of

orderon or		
Dextrin, Yellow	5	oz.
Water	3	oz.
Glycerin	5	oz.
Phenol	0.2	oz.
Ultramarine Blue Powder	75	oz.

Ultramarine Blue Paste, Laundry Glue 5 oz.

Water 10 oz. Soak cold, then warm to dissolve. Yellow Dextrin 5 oz. 3 oz. Glycerin (sp. g. 1.23) 5 oz.

Mix both parts warm, conserve with 0.2% nipagin, moldex or phenol, etc., and grind now with

Ultramarine Blue or

Imitation of Ultramarine 75 oz. formed by precipitating anilinlakes (dve-stuff) on insoluble inorganic bodies on china clay or white bolus.

Laundry Sour

U. S. Patent 1,998,819

A souring composition is formed of sodium fluosilicate 84, sodium acid fluoride 15 and gelatin 1, all parts by weight, or the like.

Cleanser for Hunting Calf Leather Trioxymethylene Cleaning Benzoline 30 cc. 5 g. Oxalic Acid Liquid Soap 20 cc. Mix thoroughly.

Cleanser for Sporting Leathers

Water	75 cc.
Acetic Acid (80%)	5 cc.
Alcohol (95%), Denatured	30 cc.

Cleaner and Disinfectant for Metal Articles

U. S. Patent 1,937,229		
Sodium Silicate (D. 1.38)	300	g.
lus 500 g. of following		
Calling II. a. 11. a. 14.		

Sodium Hypochlorite

(D. 1.125)	562 g.
Caustic Soda (D. 1.383)	250 g.
A 1/4 to 2% solution of above	is used.

Bleach-Bath for Used Oil Corks (e.g. of Olive Oil Bottles)

a. Remove fats with hot alkaline solutions, as soap, soda, trisodium phosphate; wash thoroughly with hot water.

b. Hydrogen Peroxide

(1.5-1.6%) Ammonia (25%) 200 g. Treat corks cold (18-20° C.) for about

five days, adding every 8 hours new Ammonia (25%) 40-50 g.

Oven Cleanser Formula No. 1

Olein, Distilled	40	oz.
Stearin	10	oz.
Mix warm.		
Spindle Oil	40	oz.
Tetralin	9	oz.
Ammonia (sp. g. 0.91)	1	0 £.
Emery or Pumice or Tripoli		
sufficient to make	pa	sty

No. 2

Ceresin Olein	(56–58°	C.)		g.
Mineral	Oil		17 6	g.

about 10 g.

('hromium Oxide Carborundum or Emery	15 g. about 45 g.
Printing Form and Cyli	inder Cleaner
Test Benzoline (B. P. 130-150° C.)	80 cc.
Xylol	15 cc.
Petroleum Oil	5 cc.
Ignition point should l	be over 21° C.
- ~	-

Slate Powder

Rug Cleaner

Coconut Oil Soap	12	oz.
Ammonia (28%)	2.8	oz.
Glycerin	7.9	
Water	77.3	υz.

Radiator Cleaner

Compound for use in hot	
automobile radiator flushing	tanks.
76% Flake Caustic Soda	60 lb.
Sal Soda	30 lb.
Rosin	10 lb.

Use about 40 lb. to 75 gal. water.

Dry Cleanser for Wallpapers
Wheat Starch 35 oz.
Sodium Chloride, Saturated
Solution 65 oz.

Warm upon water-bath and stir until sufficiently plastic. Shortly before the end of this treatment, when cooled, add a little naphtha. Apply like a sponge craser. Pack in air-tight tins.

Wall Cleaner

Corn Flour		90 lb.
Copper Sulphate		9 16.
Alum		1 lb.
Mix and dissolve in	boiling	water.

Scouring Soaps

The following is a soap-sand cleaning preparation that has a wide sale for household and general purposes. It takes the form of a palm oil and coconut oil soap, which is then liquored down in the same pan with carbonate of potash, carbonate of soda crystals, silicate of soda 100° Twaddell, and water.

Melt the two oils, pass in steam, and then pour in caustic soda gently, adding a little water from time to time to keep the soap smooth. Saponification will proceed fairly easily, as the palm oil soon takes up. When all the caustic soda has been added, pour in the remainder of the water in such a way that the mass never ceases to simmer; the operation should take about 4 hours. Towards the end add the other ingredients, which will dissolve easily, as the finished product is very similar to a liquid soap.

very similar to a liquid soap.

Let the soap liquid cool to about 90° F., and to 10 lb. of dried common sand add the same amount of the above soap. All the time the soap is being added, the mass must be stirred rapidly, and when it resembles a thick sludge it will be ready to pour into tims. The only precaution to take is that the mass must not be poured in too warm, as naturally the sand would precipitate in the tims; this part of the operation can only be perfected by actual experience and must always be done very carefully, but no difficulty should present itself if all directions are carried out as given.

Mixing

Coconut Oil	4	lb.
Red Palm Oil	69	lb.
Caustic Soda, 60° Twaddle	37	lb.
Additions		
Carbonate of Potash	5	lb.
Soda, Sal	15	lb.
Silicate of Soda.		
100° Twaddle	21	lb.
Cresylic Acid	3	lb.
Pine Oil	14	lb.
Orange IL (Color)	- ¥	OE.
The whole mass of soap an	d ad	ditions
hould total up to 784 lb., with	h the	addi.
ion of water.		

A hand-cleansing soft sonp can be obtained by the use of a carbolic soft soap, preferably one made from vegetable and not fish oils, using the same proportions of sonp and sand as in the previous example, but it would be better in this case to use, in place of the sand, pumice powder of 120 mesh. Sand is, naturally, coarse and cheap; better scouring agents might be used, such as silver sand, or pumice powder of 60, 90, or 120 mesh, according to the nature of the finished article desired.

Scouring Powder

Silica 100-125 mesh	75 oz.
Soda Ash	13 oz.
Trisodium Phosphate	8 oz.
Soap Powder	4 oz.

These materials in powdered form are thoroughly mixed together and are ready for use as such.

Stain Emulsifier

Liquid Soap (15%)	40 cc.
Liquid Soap (15%) Turkey Red Oil (100%)	25 cc.
Decalin	4 cc.

Turpentine	4	cc.
Ethylene Glycol	10	cc.
Methylene Glycol	10	cc.
Methanol	5	ce
Terpineol	2	CC.

Removing Glue Stains from Wood

Casein and vegetable glue stains can be almost entirely removed by sponging the stained surface with an oxalic acid solution prepared by dissolving 1 oz. of oxalic acid crystals in about 12 oz. of water. Stail better results may be obtained by moistening the wood first with a sodium sulphate solution made up in the same conventration as the oxalic acid. In this way stains have been almost eliminated.

Remover for Tobacco S	Stains on Fingers
Hard Soap Powder	40 oz.
Marble Meal	20 oz.
Alcohol, Denatured	40 oz.
Soap hands with this	s mixture, rub at

Soap hands with this mixture, rub at the same time with finest pumice powder.

Removing Pitch or Varnish from Hands or Glass

Household Scouring Powder Dutch Cleanser type

Acetone sufficient to make a thin paste

Rub the hands or article to be cleansed with this paste. The viscous impurity is at once dissolved in the acctone, and is absorbed into the powder mass. Within a minute or two the acctone evaporates, leaving a mealy or dry powder which can be dusted off, or in suitable cases as with the hands, washed off. Do not use on a painted, varished or lacquered surface, which would be injured by the acctone. This is a very economical material for the purpose.

Soot Destroyer

Salt	85	oz.
Copper Sulphate	8	οz.
Zinc Dust	7	oz.

Steamship Chimney Soap

,	<u>F</u>		
Soft Soap, Brown	20	g.	
Water	12 - 15	cc.	
Potassium Carbonate	1.5-2	g.	
Hexahydro-cresol	1.5-2	cc.	
Decahydro-naphthalene	3_4	cc.	
Sodium-Di-Isobutyl-naphtha-			
lene Sulphonate	1.5-2	g.	

Cleanser for Lampblack-Dirtied Surfaces

a. Olein or Oil Fatty Acid 45.45 kg.

b. {Caustic Soda (128-130°) 6.64 kg. Water 26.36 kg. c. Alcohol 45.4 l.

Saponify a with b on water bath, dissolve, then warm (below 70° C.) in c. Add stirring

d. Tripoli 900 kg. and thin 10 times with water.

Floor Sweeping Compound Formula No. 1

Sawdust, Dyed Green with Aniline Dye, e.g., Brilliant Green 35 kg.

Rock Salt 35-40 kg.
Mineral Oil, Deodorized
(2-3° E. at 50° C.) 25 kg.

No. 2

The following is a representative formula for floor sweeping compounds.

Tinned Ware Cleaner

Sodium carbonate alone is not a satisfactory cleanser for milk containers of tinned copper, since it slowly removes tin as stannite owing to the presence of dissolved oxygen. The exposed copper produces an "off flavor" in the milk. The addition of sodium sulphite reduces the rate of attack to nearly 0.1. It is much more effective than a number of other reducing agents tred because it is more active in reducing the amount of dissolved oxygen. Suitable proportions are 1 lb. sodium sulphite and 10 lb. washing soda, 1 lb. sodium sulphite and 4 lb. sodium hydroxide (or sodium carbonate).

Type Cleaner

Butyl "Cellosolve" Diglycol Laurate	1 pt. 1 fl. oz.

Cleanser for Velvet Shoes

Olempori Iol	1 (1100	DHOCO	
Water		100	cc.
Potassium Alum		1	g.
Alcohol		20	cc.
Turkey Red Oil		5	cc.

Composition for Cleaning Walls, Paint, etc.

French Patent 774,876

The composition contains corn flour 455, copper sulphate 40, alum 5 parts and is mixed with boiling water for use.

Painted Woodwork Cleaner

This specialty product quickly removes dirt from paint and leaves the punted surface with a bright, clean, Instrons finish. The diglycol stearate serves the combined purpose of emulsifying the dirt as fast as it is dissolved and of impunting a lasting natural luster to the cleaned surface. The product, therefore, may truly be said to both clean and shine in one operation. This new type of cleaner is made to the following formula:

Diglycol Stearato 1 lb.
Kerosene 1/4 gal.
Trisodium Phosphate 4½ oz.
Water 12 pt.

Method of manufacture: The diglycol in a double boiler until the wax is thoroughly dissolved. Kerosene is inflammable, therefore care should be taken to prevent it from catching on fire. The trisodium phosphate is dissolved in the water and heated in another container to a temperature of about 150° F. The hot water solution is then added to the hot kerosene solution while stirring at high speed. Stirring should be continued at a good rate until the mixture is of even milky consistency. Mixing may then be continued at a slow rate until the batch has cooled to around 85° F.

This product is applied in the usual manner by rubbing with a rag or cloth. The same product may also be used for cleaning automobiles before waxing. However, for this service 12 oz. of fuller's earth should be thoroughly worked into the above batch after it has cooled over night. The fuller's earth should not be added until cooling is complete. With this addition a product is produced which cleans rapidly and without scratching the finish.

"Soluble" Pine Oil Fluid

A satisfactory clear, pale straw pine concentrate, which is perfectly stable and gives a dense milky emulsion when added to water can be made from the following formula:

Heavy White Pine Oil	70 cc.
Oleic Acid	12 cc.
Water	18 cc.

The procedure is very simple—dissolve the olee and in the pine of in the cold, and neutralize carefully with a 28% solution of caustic points or soda. Caustic points gives a slightly better color than caustic soda. By this method no heat whatever is required.

Sonp Towel U. S. Patent 1,969,900

A towel for elenning surfaces consists of a paper towel currying a detergent composition including pure oil about 3-10 parts, a soap about 3-0.6 parts and water about 55-95 parts.

Sodium Metasilicate Solutions

Solutions containing 20 g. per 1, of a commercial detergent preparation (sodium sheate 40, baking soda 30, sonp-powder 20, sodium perbonte 10) show turbidity a few hours after preparation followed by precipitation; this reducts it useless. Solutions of 5-10 g. per 1, of sodium sheate begin to precipitate in presence of 35-40 g. baking soda per 1, and precipitation is instantaneous with more than 40 g.; a solution of 15-30 g, per 1, of sodium silicate begins precipitating in presence of 20-25 g. baking soda. Substitution of trisodium phosphate for laking soda immediately corrects the

Movie Film Cleaner

MOVIE PHILI ORALIEI		
Carbon Tetrachloride	65	oz.
Ethylene Dichlorido	10	oz.
Petroleum Ether	25	oz.
This composition is used to	clean	dir

This composition is used to clean dirt, greasy spots and all foreign matter off of both faces of a movie film without affecting or having any solvent action on the film or gelatin coating itself.

The petroleum ether is a light fraction distillate with an end point under 100° C. These solvents are mixed together and are then ready for use.

Benzine Soap

Dissolve 10 lb. of card soap in boiling water, add a strong solution of magnesum sulphate slowly with stirring until it is all transformed into an insoluble mass, skim off the magnesium soap thus formed and purify by boiling it with fresh water. Remove the excess of moisture by squeezing through a cloth and pressing. Place the soap in a jacketed copper kettle and heat slowly to 266° F., turn off the heat and add 7 lb. of decodorized petroleum distillate. Dissolve

the product in 22 gal. benzine. If the solution is not clear the water has not been completely removed. For garment cleaning use 1 qt. of this solution for 25 gal, of benzine.

Dry Cleaning Solvents for "Celanese"

The following chemicals are safe for cellulose acetate fabrics: gasoline, Stod-dard's solvent, cleaner's naphtha, kerosene, dilute alkalies (such as soap and water, soda, ammonia, sodium hypochlorites, Javelle water and washing sodas), glycerin, carbon disulphide, turpentine, all the hydrosulphite solutions (such as decolorite, blanket, sulphogen, burmol, paragene and lykopon), petroleum ether, vaseline, toluol, xylol, good grades of wood or denatured alcohol used cold and washed thoroughly, sulphuric ether, tri-chloroethylene, benzol, which is one of the best all around spotting chemicals, and unadulterated carbon tetrachloride, which is rapidly taking the place of chloroform. It is a known fact that carbon tetrachloride will absorb a small amount of moisture from the air if the container is left open. If moisture is present this powerful solvent is crippled and will not be as effective as when dry. To test carbon tetrachloride for purity, take two parts mineral oil, such as Nujol, and one part carbon tetrachloride. Mix. If this mixture becomes milky it denotes the presence of water in the carbon tetrachloride and in this condition should not be used for spotting purposes.

Dry Cleaning Soap

Curd Soap		30	oz.
Water		40	oz.
Ox Gall (Dried)		10	oz.
Soda Ash		5	oz.

Shred the soap and dissolve in hot water, adding the ox gall and soda. Evaporate the solution until on cooling, a sample on a slab sets solid. Pour the mixture into trays or molds. The disadvantage of such a preparation is its rather unpleasant smell.

Dry Cleaning Soap British Patent 407,088

Fourteen and two-tenths grams of sodium hydroxide is dissolved in 25 cc. water and stirred into 100 g. oleic acid and 100 cc. trichloroethylene; 70 g. tri-ethylene glycol or 50 cc. diethylene glycol is added and the product is dissolved in trichloroethylene for dry cleaning.

Textile Soap French Patent 658,412

Castile Soap	200 lb.
Tallow Soap, Powdered	95 lb.
Soda Ash	20 lb.
Borax	10 lb.
Turpentine	25 lb.
Caustic Alkali	20 lb.
dissolved in 30	

Trian 900

Arer Soup	
Red Oil	2050 lb.
Rosin	1050 lb.
Soda Ash	290 lb.
Caustic Soda (50° Bé.)	746 lb.
Water to make	11000 lb.

Ox Gall Soap

Since ox gall derived from bile has an unpleasant smell, an improved method is to add to soap solution about 4% of sodium cholate, the sodium salt of choice acid which is a purified decomposition product of bile. It is claimed thus that the advantages of the detergent power of ox gall are obtained without the accompanying odor.

Rose Sonn

	White Tallow Scap Moistened Cinnabar	10000 kg. 60-80 kg.
	Rose Essence Clove Essence	40 kg. 15 kg.
с.	Cinnamon Essence Neroli Essence Bergamot Essence	10 kg.
	Neroli Essence	10 kg.
	Bergamot Essence	30 kg.
	Perfume	

Windsor Soap

10000 kg.

a. White Tallow Soap

	Bergamot Essence Caraway Essence Clove Essence Thyme Essence	60	kg.
b	Caraway Essence	25	kg.
	Clove Essence	16	kg.
	Thyme Essence		kg.
	Perfume		
	or		

- 1	Bergamot Essence	25	kg
٠ ا	Caraway Essence Rosemary Essence	60	kg.
0.	Rosemary Essence	15	kg.
	Fine Lavender Essence		kg.
-	Fine Lavender Essence	15	1

Witch Hazel Soap

Witch Hazel Extract U.S.P.		oz.
Distilled Water	10	0 z .
Triethanolaminelauryl-		
enlphonate	80	OS.

Perspiration Odor Destroying Soap Aluminum Chloride Crystals Hydrochloric Acid 1/10

Normal Tricthanolaminelaurylsulphonate

1-2 oz. 96-95 oz.

Soft Soap Manufacture

Soft soap contains normally 40 to 44%. of fatty acids. The best method of saponification is to take the calculated quantities of alkali sufficient to effect complete saponification, with an excess of 1 to 1.5% alkali. The caustic solution, preferably of a density of 30° Tw. (about 19° Bé.), is brought to a boil and the melted charge added as quickly as possible without the contents frothing over. Emulsification follows with rapid suponification. The process is usually complete in a few hours' time, water being added when necessary. If rosin is to be incorporated, it is best added after the other stocks have been saponified.

Unless caster oil is present a soft soap charge cannot be worked with caustic soda alone. With caustic soda, castor oil will form a soft soap. Soft soaps can be made with castor oil in which varying proportions of other stocks have been introduced with the substitution of varying proportions of caustic potash for caustic soda. Saturated fatty acids tend to give stringy soap even with potash. higher these are in the homologous series, the more pronounced is the stringmess.

The percentage of caustic soda which can be substituted for eaustic potash will depend on the percentage of castor oil introduced into the blend. Practical experiments indicate that about 2% of caustic soda can be substituted for emistic potash for every 1% of castor oil introduced into the blend. Use of caustic soda in this way does not affect translucency and gloss.

Linseed-soda soap is stringy, but the corresponding potash soap is non stringy with a desirable body. Peanut oil soda soap is stringy but the potash soap is not. The following blends suggest the possibilities for soft soap manufacture:

The charges given below produce a stringy soup with 80% caustic potash and 20% equivalent caustic soda, but have the right non-stringy body with caustic potash only:

Formula No. 1

Peanut oil 20 parts, linseed oil 50, cottonseed oil 20, tallow 5 and rosin 5.

No. 2

Peanut oil 20 parts, linseed oil 50, cottonseed oil 20 and tallow 10.

No. 3

Linseed oil 60 parts, cottonseed oil 30 and rosin 10.

No. 4

Linseed oil 65 parts, cottonseed oil 25, rosin 10.

No. 5

Linseed oil 67 parts, peanut oil 13, cottonseed oil 10, tallow 5 and rosin 5 can be used with 80% of caustic potash and 20% equivalent caustic soda to give an almost non-stringy soap with only a slight thready tendency.

Linseed oil 73 parts, cottonseed oil 15, rosm 10 and coconut oil 2 can be used with 70% of caustic potash and 30% equivalent caustic soda to give a non-stringy soap. In general it is preferable to use more potash. This represents the lower limit of potash with this type of blend.

No. 7

Linseed oil 73, castor oil 20, rosin 5 and coconut oil 2 gives a correct nonstringy soap with 60% caustic potash and 40% equivalent caustic soda, due to the introduction of castor oil.

The following blends with higher percentages of easter oil give non-stringy soap with caustic soda alone:

No. 8

Linseed oil 38, castor oil 50, coconut oil 2 and rosin 10 parts.

No 9

Linseed oil 32, caster oil 45, coconut oil 3 and rosin 20 parts.

No. 10

Linseed oil 50, castor oil 35, coconut oil 3 and rosin 12 parts.

Soap Rancidity, Preventing

This is best done by kneading into the dry soap, before milling, .7% of the following mixture:

Beeswax 300, anhydrous lanolin 400, liquid paraffin 390, water 300, borax 17, sodium thiosulphate 690, water 200. Melt together the wax, lanolin and paraffin oil; then dissolve the borax in 300 parts of water and pour this solution in a thin jet into the hot mass of malten fats at a temperature of about 95° C. Boil for a few minutes longer, then set uside and let cool to 50°, stirring frequently. Pour the hot solution of sodium thiosulphate in 200 g. of water into the fat-borax emulsion in a thin jet and stir until smooth. In some cases, for example en using an unusually large quantity of perfume, it is advisable to add 1% of

the following:

Beeswax 200, anhydrous lanolin 600, liquid paraffin 390, water 200, borax 17, sodium thiosulphate 690, water 200, sodium silicate 450, granulated sugar

Superfatting Soap

Use of a superfatting agent undoubtedly improves the texture of soap, making it more plastic and easily worked. It also tends to neutralize any alkali which might be present, and thus remove harshness which might irritate sensitive skins. A good mixture for this purpose consists of equal parts of stearin and white petroleum jelly, or 2 parts stearin, 1 part lanolin, and 1 part white petroleum jelly. These are melted, mixed, allowed to cool, and 1 to 11/2 lb, added per 100 lb. of chips added with the other ingredients at the mixing stage.

Soap Spirit

Olive Oil 1000 cc. Caustic Potash (50%) about 396 cc. Distilled Water 2600 сс. Alcohol (90%) 6000 cc.

Softener for Hard Water

Water Glass (36-38° Bé.) 25 oz. Water 25 oz. Ammonium Carbonate about 50 oz. Mix well (warming), pour off to solidify the paste. When cool, grind and add to 95 oz. of the material. Trisodium Phosphate 50 oz.

TEXTILES, FIBERS

Starches and Sizes for Cot	ton Sheeting	1 Gum Arabic	8 g.
Formula No. 1		Soap	32 g.
Cornstarch	100 lb.	Glycol Stearate	18 g.
Castor Oil	⅓2 pt.	Bornx	2 g.
Color	to suit	Pepsin	0.15 g.
	0-240 gal.	No. 2	
	.,	Lauric Acid	20 g.
Boil together until smootl	1.	Gelatin	34 g.
No. 2	100 11	Soap	30 g.
Cornstarch	100 lb.	Gum Arabic	8 g.
Gypsum	80 lb.	Ethylene Glycol	6 g.
Castor Oil	1 pt.	Borax	2 g.
Color	to suit	Trypsm	.05 g.
	0-240 gal.	Olue No. 3	20 g.
No. 3	60 lb.	Gum Arabic	
Cornstarch	5 lb.		_ B.
Lard	2-4 oz.	Glycol Stearato	
Blue Dye		Soap	
Water	120 gal.	Glycerin Borax	
No. 4	es 11.	Pepsin	
Cornstarch	65 lb. 10 lb.	1 epsin	.01 g.
China Clay	5 lb.		
Laid		Rayon Sizo	
Color Water	to suit	Calcium Resinate	20 lb.
	120 gal.	No. 1 Lard Oil	10 lb.
No. 5	!!	Xylol	35 lb.
Potato Starch	100 lb.	Damar Gum	10 lb.
Steeped Flour (24° Tw.)	10 gal.	Mampulation: Dissolve the	damar gum
Slaked Lime	15 gal.	in the xylol and add the other	
China Clay	15 gal.	at 50° C. Then cool slowly	
Elaine (Red) Oil	3 pt.	tion.	-
Blue Color	12 oz.		
Water to make	120 gal.	Light Goods Sizin	•
Boil for 1-2 minutes.			
-		Formula No. 1	
Cream Sizing		Soluble Potato Starch 11/2	
Tallow	36 lb.		-5 pt.
Calcium Chloride	6 lb.	Water	5 gal.
Starch	7 lb.	The starch and glucose	are entered
Gum Arabic	6 lb.	into the water and the whole	
Water	45 lb.	a boil and continued at that	
Cook and stir at 220-230°	F. for 1-2	until the starch particles	are entirely
hours.		cooked, which will depend up	
		ticular type of starch us	
Sizing Rayon and Silk		using, the mixture should be	allowed to
		cool to a temperature of ab	
French Patent 779,584		The purpose of the glucose	
Rayon and silk are sized to give firm-		a soft feel to the material	and may be
ness, elasticity and suppleness by a solu-		omitted.	
tion in water of		No. 2	
Formula No. 1		Another mixture that is	
		setting goods other than	
Stearic Acid	15 g.	structed of rayon, is to 1	
Glue	35 g.	white finishing gum to 5 gal	. water.

A mixture	that 1	nay be	recomm	ended
for producin	g a sof	t, lustro	us finis	h, and
particularly	for ra	yon bra	.ids, 18	given
below:				

a. Gum Arabic dissolved in 1

gal. Water Gum Tragacanth dissolved in 1 gal. Water

Use one part solution a and one part solution b to 4 to 5 parts water and apply lukewarm.

Running the goods through plain, lukewarm water and then through the calender will often remove wrinkles that have been developed in the process of dyeing.

Glue is the substance most often employed for stiffening braids, as well as other textile fabrics. This ingredient comes in many different qualities, and the grade required will depend upon the quality of the material to be treated and the result desired. The flakes or granules of glue should be allowed to dissolve in water some time before it is to be needed at the finishing machine, and as glue varies greatly, it is advisable to experiment with each new lot before proceeding with any quantity of material.

Various substances are used to prevent the size bath from souring. Among these are zinc chloride, sodium fluoride, bluestone, and formaldehyde. Any of these chemicals are used in very small quanti-

ties.

Textile Size

Glucose	7 lb.
Soluble Oil	3 lb.
Magnesium Sulphate	1 lb.

Textile Paste or Size

Potato Starch	100 lb.
Calcium Chloride	300 lb.
Water	300 lb.

Manipulation: Soak starch and calcium chloride in the cold water for 2 hours then gradually heat mixture to boiling. Boil for 1 or 2 hours until a thick paste is formed.

Equipment: Clean wooden vat with open steam for boiling.

Cleaning Solvents for Textiles Formula No. 1

Carbon Tetrachloride

No. 2

Carbon	Tetracblo	ride	850	cc.
Heavy	Benzoline,	Purified	150	cc.

No. 3	
Heavy Benzoline, Purified	640 cc.
Ethyl Ether	120 cc.
Turpentine Oil, Purified	120 cc.
Ethyl Acetate	120 cc.
(Inflammable!)	
No. 4	
Heavy Benzoline, Purified	600 cc.
Turpentine Oil, Purified	120 cc.
Ethyl Ether	160 cc.
Ethyl Acetato	120 cc.
(Inflammable!)	
No. 5	
Carbon Tetrachloride	650 cc.
Alcohol	100 cc.
Ethyl Ether	100 cc.
Heavy Benzoline, Purified	80 cc.
Soap Spirit	50 cc.
No. 6	
Trichloroethylene	

Scouring Rayon Circular Knit Fabric 1. Run water in kettle (80-120° F.) using minimum amount that will enable

- the fabric to run freely over the reels. A properly loaded kettle of the correct type requires approximately a 20 to 1 bath.
 - 2. Load kettle with fabric.
- 3. Add 2 lb. soda ash or trisodium phosphate (depending upon water conditions).
- 4. Turn on steam and run goods for 10 minutes.
- 5. Add 3 lb. high grade neutral soapolive or red oil base.
- 6. Add 2 lb. "soluble pine oil" or a similar solvent containing material. If desired this solvent material and soap can be added simultaneously in order to aid solvent dispersion.
- 7. Raise bath to boil. Observe condition of bath at all times. If bath does not show a good, clean, sudsy condition, add more soda soap and pine oil. It is impossible to accurately predict the amount of soda soap and solvent or the exact proportions of the same that will be required under an unknown set of condi-
- 8. Run the kettle at or near the boil for 1 hour.
- 9. Drop bath and proceed with bleaching or dyeing operation.

Cleaning Tent Canvas

Mildew can be removed from a tent by sponging the canvas with a weak solution of calcium hypochlorite, or bleaching powder. Be sure to wash the solution out well after using.

Cotton Textile Printing

For the shading of the pink print a paste is prepared with 40 parts of Irisamine G, that are dissolved in 400

parts of iron-free water. The resulting solution is then incorporated into 500 parts of starch tragacantl thickening, warming for a short time, agitating until the mass reaches 60-70° C., and entering 80 parts of acetate of chrome at 18° Re, and bringing to 1000 parts through adding more water if this is necessary.

The starch tragacanth thickening, required in the above case, is prepared with 60 parts of wheat starch and 50 parts of wheat flour, that are made into a uniform semi-transparent paste with 700 parts of water, adding to this while still boiling 200 parts of a 6½% gum dragon muciage and 30 parts of olive oil. The bath being brought with water to 1000 parts

in all.

For the back of the pink print 200 parts of the above color paste are measured out and mixed first in a warm bath containing 800 parts of the starch tragacanth thickening, and then with 2 parts of acetate of chrome at 18° Bé. and 5 parts of acetic acid at 6° Bé., that are added at the right moment in the cooling down bath.

The shading product, needed for the red print, is obtained with 10 parts of a suitable brand of safranine, that are dissolved in 90 parts of acetic acid at 6° B6., 10 parts of acetin and 300 parts of iron-free water. The resulting solution is then added into 500 parts of the starch tragacanth thickening indicated above, and after cooling sufficiently (60-70° C.) are entered 60 parts of a 50% tannin acetic acid solution and 40 parts of acetic acid at 6° B6., bringing the whole to 1000 parts with further water.

For the backing of the red print a fourth printing paste is prepared with parts of a suitable safranine, dissolved in 20 parts of acetic acid at 6° Bé., and 100 parts of inco-free water. The resulting solution is then poured into 550 parts of the starch and gum dragon thickening, and when this has been properly incorporated, steam is turned off, and the bath is left under the action of the agitator until 70° C. has been reached. Fourteen parts of a 50% tannin acetic acid solution, and 20 parts of acetic acid at 6° Bé. are then poured in, in close successions.

sion. The bath is made up after this to 1000 parts with further water.

The cotton cloth is printed with the above four color pastes, dried by passing through the hot-flue, and steamed for 1 hour without pressure, or for half this time with one half atmosphere. After the steaming, the goods are treated in a 1% tartar emetic bath at 50° C, rinsed for some time and dried. If the free acid in the goods is not eliminated in this way, the cotton choice is passed through a second bath containing from 5 to 10 parts of chalk per 1. of iron-free water, giving a second rinsing, and drying and finishing.

If the printing is to be conducted on a pure white cotton cloth, the cost of treatment is much reduced, as a direct printing process is only required. This can be conducted with one of the pastes given below, the first of which requires, after its application and drying, a two hour steaming at 1 atmosphere pressure, while the second needs instead a one hour steaming with one-half atmosphere. Both colors being improved by a soaping.

Formula No. 1

Two and a half parts of alizarine black in paste S are mixed with ½ part of a sectic acid at 6° B6, and 6½ parts of a suitable starch thickening. The mixture is warmed until obtaining uniformity. After this it is allowed to cool down somewhat, and is entered ½ part of acctate of chrome at 20° B6, bringing to 10 parts in all with water.

No. 2

Three hundred parts of a suitable brand of chrome orange are incorporated with 620 parts of acid thekening and 80 parts of acid thekening and 80 parts of acid thekening and 80 parts of acid. The acid thickening is prepared by boiling 210 parts of wheat starch with 570 parts of iron-free water, after having conducted properly the mixing in the cold. When a semi-transparent adhesive has thus been produced steam is turned off, and toward 70° C. are entered 220 parts of acetic acid at 6° B6., bringing with water to 1000 parts in all.

If the cotton material is colored in a

If the cotton material is colored in a light pink, this is obtained by dyeing on the jigger or on the padding machine with a suitable bath of Erika GN, shaded or not with Chrysophenine G; with a hath or Benzo fast scarlet 4BS (using the correct percentage), of Diamine rose BD, or of any other substantive pink; rinsing, drying and printing with the following

color paste:

Seventy-three parts of Ciba red G in paste are mixed with 27 parts of a 33% British gum thickening, and passed through a fine sieve, bringing then with water to 100 parts. Fifty-five parts of the above mixture are then entered in 12 parts of further 33% of British gum thickening, adding a little later 20 parts of caustic soda lye at 36° Bé., and 6 parts of glycerine. The whole is warmed just sufficiently for obtaining a uniform incorporation, and after having allowed the bath to cool down to about 50° C, are entered 7½ parts of hydrosulphite NF concentrated, bringing with water to 100 parts.

When the cotton cloth goods have been printed with the above pink paste for obtaining the necessary details in the flowers and in the dark ground, the material is dried and steamed from 4 to 5 minutes at 105-107° C., being then left to hung for a short time, and finally treated with a bath furnished with 5 parts of olive oil or cottonseed oil soap and 2 parts of calcined earbonate of soda for every thousand parts of iron-free water, the bath being kept all through towards 60° C. After this the goods are given a last drying and are finished.

Logwood Speck Dye Logwood Extract 51° Tw.

Logwood Extract 51° Tw. 48 lb.
Soda Ash 30 lb.
Bluestone 12 lb.
This should be diluted to about 2-3°

Garl D.

Scal Brown Cotton Dye

Cutch 35 lb.
Hypernic Extract 16 lb.
Logwood Extract 31/2 lb.

Add to dye bath and boil until dissolved, then add 3 lb. bluestone, add edd water, rake well and enter yarn. Give 6 turns and put down over night. Take up, give 6 turns, introduce into a solution of 4 lb. chrome at 160° F. and give 6 hours. Remove, wash well in cold water, put back in cutch liquor, 6 turns; into chrome, 4 turns; into chrome, 4 turns. Wash off each time after chrome. Start new kettle with

Fustic Extract 7 lb.
Logwood Extract 31/2 lb.
Boil well for 2 hours.

 Violet Logwood Textile Ink

 Logwood Extract (Weak)
 300 lb.

 Alum
 12 lb.

 Dextrin
 15 lb.

Dissolve the alum by heating in a part of the extract solution. Finally 1½ lb, finely powdered lead acetate are slowly added and dissolved.

Textile Padding Liquor

Acetic Acid 50% 3 gal.
Formic Acid 85% 3 gal.
Glauber's Salt Crystals 40 lb.
Water to make 100 gal.

The goods are padded on the face and the drying cylinders must not be too hot at first, so that sticking of the prints may not take place. Moderate drying in the initial stages should be the rule but at the same time if duying is not carried out properly there will be a grave danger of marking-off on the cylinders if any of the print color is allowed to adhere during the process. The wrapping of the first cylinder is sometimes advised in order to prevent sticking, but the circumstances in each case will dictate the precautions which will have to be taken. Two or three cylinders in any event will be found sufficient for the full development of the colors.

Preparation of Print Colors

In using the powder brands the following method of producing a print color is normally adopted.

Dyestuff Powder 8-16 oz. is pasted with Caustic Soda Solution

(70° Tw.)

Monopol Oil or Similar

Soluble Oil

45-14 pt.

15 pt.

Neutral Chromate Solution ¼ pt.
The mixture is then allowed to stand for a short time before being added to

Water 2-3 pt.

Starch-Tragacanth (Thickening as Required)

Making the whole up to

Printing Paste 1

For the production of lighter shades from the above standard a thickening of the following type is made up.

Neutral Starch-Tragacanth 1 gal. Caustic Soda Solution (70° Tw.) 44 pt.

Neutral Chromate Solution 4 pt.

The neutral chromate solution is prepared in the following manner:

Sodium Bi-Chromate Crystals 1½ lb.

dissolved in Water To this add	6	pt.
Caustic Soda Solution (70° Tw.)	22	oz.
Make up to	_	
Neutral Chromate Solution		gal
The paste brand dyestuffs as follows:	tre pi	epare
Dyestuff	1	pt.
Neutral Chromate Solution	8	oz.
Monopol Oil	1/1	pt.
Water	2	pt.
Neutral Starch Tragacanth	5	pt.
Printing Color	1	gul.
Thickening for Hand Printi Formula No. 1	ng or	Sılk
Mix White Starch	5	lb.
and	•	•17•
White Dextrin	5	lb.
Acetic Acid, 12° Tw. Olive Oil	71/ <u>2</u> 2	lb. lb.
and then add	-	•••
Water	016	gul.
Boil to a paste.	- /2	B
No. 2		
Mix		
White Starch	5	lb.
with Water	1	gal.
and		
Glue	21/2	lb.
previously dissolved in		
Water	21/2	gal.
Boil to a paste, cool and add		
Acetic Acid, 7° Tw.	5	lb.
Olive Oil Stir well.	2	lb.

Coloring Bone Articles

The chief difficulty encountered in coloring bone material such as chess and other game counters, buttons, horn handles for umbrellas and walking sticks, ornamental vases and similar brica-brac of this type, etc., consists in obtaining good penetration of the dye. It is an unfortunate fact that certain acid and hasic dyes of poor fastness to light will penetrate bone material better than some of the faster colors. Where penetration is too shallow, bone articles subjected to much handling like chess and draughtsmen, umbrella and walking stick handles and so on, soon disclose unsightly light

places where the superficial film of coloring matter has worn off.

Bone material is commonly dyed in a nested copper kettle, the inner container which carries the stock being perforated with small holes for the circulation of the liquor. The container can be lifted from the outer easing when it is desired to examine the stock during processing. Coloring of bone material is usually performed before it is polished, as treatment in hot liquor would roughen the surface of polished goods. When small articles like buttons, electric bell and light switch press plungers, ivory sectors for inlay and unriqueterie designs and so forth are to be colored, handling of the stock is facilitated by processing it in bags of linen net, each bag having a capacity of about S oz. of stock.

It is customary to boil-off bone material in clean water before coloring it. If the stock contains traces of oil or grease acquired during turning and fret-cutting of ornamental pieces, a small amount of pearl ash is put into the boil-off bath in order to emulsify the fatty substance. It is well to be sparing in the use of the alkah because the employment of an excessive amount will turn the bone a yellowish color. The use of soap for boiling off is also apt to bring about this yellow discoloration in the stock; moreover, the presence of residual soap during coloring of the bone material will hinder penetration. The usual duration of the boil off is from 15 to 60 minutes, according to the size of the pieces in the stock and the kind of bone. Antler and tusk material is harder and less porous than stock manufactured from sawn bone of bovine origin.

When the stock has been taken out of the boil off kettle, it is plunged into the boiling dyclath, which is already fully charged with the appropriate dycatum. Boiling proceeds for 30 to 60 minutes and then the stock is allowed to steep in the cooling bath for several hours in order to encourage penetration. It is not always advisable to process thin pieces made of horn at boiling temperature for longer than a few minutes, because of the risk of distorted material through softening of the structure in the hot liquor.

The following dyes may be employed for processing first-to-light colors on bone material. Afterohrome Black of the PV type; Alizarine Brilliant Green G; Cloth Fast Yellow R; Eriochrome Red G; Erio Fast Brilliant Blue 3R; Radio Brown B; Cutch Extract; Logwood Extract. After-chrome Black is applied to bone material in a boiling bath containing 1% of 30%

acetic acid. After processing for half an hour, 1% of sulphuric acid 168° Tw. is added and boiling is continued for a further half hour. The stock is then allowed to steep in the cooling bath for some hours, after which it is plunged into a fresh bath containing a boiling solution of bichromate of potash, the amount employed being from 1 to 2%. After 15 minutes processing at the boil, steam is turned off and the stock is left to steep for a further period of 15 minutes and then it is lifted and rinsed in warm water.

Alizarine Brilliant Green G yields fine blue-green hues of high fastness to light on clean white bone stock; when this solor is used on discolored stock, or the darker sorts of horn material, the shade which ensues is a bottle-green color.

Alizarine Brilliant Green G has good affinity for bone when applied in a boiling neutral bath. For deep shades with this dyestuff, an addition of 1% of acetic acid should be made to the bath after cessing neutral for half an hour. Cloth Fast Yellow R also possesses good affinity for bone in neutral liquor. Deep hues may be processed with an addition of acetic acid, this to be put in when the bath has boiled for half an hour. Eriochrome Red G yields rich red on bone stock. Dyeing should be commenced with the addition of 1% of acetic acid and when the bath has boiled for half an hour, 1 to 2% of bichrome may be put in. If the stock is hard tusk, boiling should be kept up for an hour before the bi-chrome is used. Erio Fast Brilliant Blue 3R produces a lively and very durable reddish-violet color on clean white bone material. This dyestuff has very good affinity for bone in a neutral bath. processing a full shade, an addition of of acetic acid may be made after the ath has boiled one hour.

Radio Brown B is a useful dyestuff for processing light or dark brown hues of first-rate fastness to light on bone stock. The affinity in a neutral bath is not good, hence an addition of acetic acid may be used at the commencement of dyeing. After the bath has been boiled for about half an hour, the color may be exhausted by an addition of 1% of sulphuric acid.

Cutch extract is an old favorite amongst bone dyes. This substance yields olive-gray to rich brown huse on bone, the shade depending on the processing method adopted. To produce olive-gray on bone stock, the material is boiled for 30 minutes in a bath containing 10 to 20% of dry cutch extract, and 1-2% of assetic acid. Steam is then cut off and

the stock allowed to feed in the cooling bath for 8 to 10 hours. The material is then put into a net bag and suspended in an empty barrel into which steam is blown for 10 minutes. The jet of steam must not impinge directly upon the stock. Oxidation of the cutch which has been absorbed is then completed by exposing the bone pieces to the air while they are spread out in shallow trays. In order to develop the olive-gray coloration, the stock is plunged into a boiling bath containing 2 to 5% of green copperas. Steam is cut off after the material has boiled for 15 minutes, after which the stock is left to steep for half an hour and then rinsed. The olive gray hue produced in this manner has long been a popular color for the bone platings on pocket knives. If it is desired to process orangebrown or deep reddish brown with cutch. development of the color is done with bichrome and copper sulphate instead of green copperas. When deep colors are green copperas. When deep colors are being processed on bone material with cutch, or other natural coloring matters, it is usually necessary to remove the film of loose color and resinous impurities which forms on the surface of the bone during processing. If this film is not cleaned off, it clogs in the bone and hinders development of the final color during after-treatment with the metallic salts. In order to cleanse the stock, the pieces are put in the loose condition into a tumbler apparatus containing a thin paste of sawdust and water, or preferably cow dung and water. When the device is set into motion, the movement of the stock in contact with the sawdust, etc., cleanses away the film.

Logwood extract is sometimes combined with cutch for the purpose of modifying the tone of the latter. Logwood extract is also used for deep black on bone articles, the process consisting in boiling the stock in a solution of logwood extract, followed by the oxidation of the hematine by steaming and exposure to the atmosphere. After the material has been freed from film in the tumbler apparatus, the black color is developed in a boiling bath containing copper suphate and green copperas. A black of this kind is not as fast to light as afterchrome black, but penetration is frequently better than in the other instance.

Dycing Vegetable Ivory Buttons

The following is suggested with the use of basic dyes: The buttons are boiled in water for 1-2 hours before dyeing. Pale shades are dyed for 2 hours at the boil in

a neutral bath; if the water is very calcareous, some acetic acid must be added.

Full shades are first mordanted for 4 hours in a bath prepared with 40 parts tannin per 1000, then rinsed in cold water and treated in a bath prepared with 20 parts tartar emetic per 1000 for ½ hour at 120-140° F. The buttons are then rinsed for ½ hour with boiling water in order to remove the free mordant and dyed in a fresh bath acidified with acetic acid.

Dyeing Brush Bristles

When dyeing fiber materials to be used for the manufacture of brushes, etc., and necessitating the material being dyed through well, it is best to use a combination of about 2-3% of a direct black and 2-4% logwood extract.

Charge the starting bath with 2% ammonia and ¼-½% soda ash, add 2-3% dye previously well dissolved in condensed water, and then about 5% cryst. Glauber's salt; boil up well, enter the material, work for 5-10 minutes, cover with a lattice frame weighted with stones, boil for 2-3 hours, and allow to feed for ½-1 hour in the cooling bath. Then lift the material, allow it to lie exposed to the air for several hours, and enter into a fresh bath heated to 30-40° C. (85-105° F.) containing pyrolignite of iron of 4-7° Tw.; leave in this bath for ½-1 hour, throw out and leave exposed to the air for several hours, rinse well and dry.

If so-called patent or luster-fiber is to be produced, the method of working is exactly as described above; only the fiber is finally taken through a bath of 40-50° C. (105-120° F.) charged as follows:

Liquor		10	gal.
Gelatin Glue		2	lb.
Soft Soap		2	lb.
Logwood Extract	;	2	lb.
Fustic Extract			
Pyrolignite of I	ron	1/2	lb.

Treat the goods in this bath for 30 minutes, allow to drain, and brush dry with suitable brushing machines. If the fiber is not lustered, 8 oz. of whitening per 10 gal. liquor are added to the bath of pyrolignite of iron.

The dye liquors may be used repeat-

The dye liquors may be used repeatedly; dyeing in the standing bath requires about ½-% of the stated quantities of dye and logwood extract, equal quantities of soda and ammonis, and about 3% salt calculated on the weight of the goods.

Coconut Fiber Dyeing
Dyestuff 30 lb.
Acetic Acid, 30% 90 lb.
Glycerin (only where the goods will be steamed after printing) 30 lb.

Water 400 lb.
Tragacanth Thickening 450 lb.

If the mats are to be steamed, the operation is carried out in a cottage steamer, the duration of steaming being from a quarter to half an hour without pressure. The mats are hung on rustless metal hooks riveted into movable metal strips which span the interior of the steaming cottage. The stock is seldom washed after steaming, unless the thickening has been made too good with the result that the printed portions handle stiffly. Basic dyes are apt to lose depth during washing, even when the stack has been steamed; hence, washing is only done where the necessities of the case call for it. Some printers regularly make an addition of tunnic acetic acid to the print color in order to heighten the real tance of basic color to washing and to general wear in the domestic sphere. The following basic colors are suitable for use in printing coir matting: Phosphine, rhodamine, magenta, safranine, methylene blue, malachite green, methyl violet, bismarck brown, jute black.

Substantive dyes prove useful for printing coir in designs of good fastness to washing. This class of dyes should be steamed after printing in order to obtain good results. The printing paste is made as follows:

3 lollows:			
Substantive Dyestuff	30	lb.	
Water	370	lb.	
Phosphate of Soda	30	lb.	
Glycerin	70	lb.	
Tragacanth Thickening			
(40 - 1000)	500	lh.	z

The following substantive colors are suitable for printing coir: Chrysophenia G, Direct Fast Scarlet 4B8, Benzopurpurine 4B, Direct Bordeaux 6B8, Direct Brown G, Direct Brown M, Direct Fast Pink BK, Direct Green B, Direct Sky Blue FF, Direct Black BH, R, E. After the mats have been printed, they are allowed to become partially dry and then they are steamed without pressure for half an hour. They are then rinsed in cold water.

Bleaching Coconut (Coir) Fiber

The bleaching process with hypochlorite is carried out in a cold bath after the coir stock has been boiled out in a solution of caustic soda. From 3 to 7 lb. of

commercial hypochlorite of soda solution are used per 100 gal. of water in the bleach bath. The stock is allowed to remain in the kettle for from 1 to 8 hours after which it is soured in a fresh, cold bath containing 1½ pt. of hydrochloric acid, 30 to 34° Tw. per 100 gal. of water, and subsequently well rinsed. The batch is then ready for antichloring, this process consisting of immersing the corf or a period of 10 minutes in a fresh, cold bath charged with 1¼ lb. hyposulphite of soda crystals per 100 gal. water. After this has been done, the stock is thoroughly rinsed in cold water, then steeped for several hours in two or three changes of water and finally centrifuged.

To bleach coir stock with permanganate of potash and bisulphite of soda, the material is first boiled out in a kettle with 3% caustic soda and after being rinsed, it is immersed for 12 hours in a cold solution of permanganate of potash, % Tw. The stock is then rinsed and entered into a fresh cold bath containing solution of bisulphite of soda % Tw. When the stock has steeped for one hour, the bath is let down, the material being then given two cold rinses. If it is then found that decolorization is insufficient, the operations just outlined are repeated.

In a case where hydrosulphite is chosen as the decolorizing agent, the stock is first soaked in cold water for 24 hours to remove the looser class of impurities and then a liquor containing 10 to 15 lb. of hydrosulphite per 100 gal. of water is prepared in a separate kettle connected by piping to the other one. The solution of hydrosulphite is then run in at a temperature of about 85° F., circulation of the liquor being kept up for 20 minutes or so by means of a rotary pump attached to the apparatus. After this period has plapsed, steam is turned on and the kettle raised to about 170° F. and maintained at this temperature for from 1 to 4 hours. If the stock is heavily colored with natural pigment, further amounts of hydrosulphite are added to the kettle from time to time. When decolorization is deemed sufficient, the bath is let down and the stock is well rinsed in cold water.

Some manufacturers of coir mats prefer to decolorize the stock in the woven
condition. In this event, the mats are
either strung on rods which rest upon the
rim of the kettle or else they are
processed in a package apparatus. This
is of an extremely simple type, it consisting of little more than an open kettle
fitted with a rotary pump for circulation
purposes. It is customary to place a
wooden trammel or grid on top of the

pack to circumvent floating of the stock due to the formation of steam pockets.

Bleaching Vegetable Fibers German Patent 615,680

Steep for 10 minutes in hot water and then place in bath containing 2.2 g. active chlorine and 1.5 g. caustic soda per l. at 32° C. Raise temperature to 75° C. and treat with hydrogen peroxide, then rinse.

Bleaching Mohair Cotton Fabric

The cloth, which is first thoroughly scoured in a soap soda ash bath, is transferred to a winch containing 500 gal, of water at 100° F. Five lb. of potassium permanganate carefully dissolved in lukewarm water are slowly added through a fine sieve. The cloth is run in this bath for 11/2 hours. After two cold 10-minute rinses the box is filled to the same height as before with cold water and 4 gal, of 72° Tw. sodium bisulphite liquor are added. The cloth is run several minutes before adding 12 lb. of commercial sulphuric acid previously diluted by pouring into several times its volume of cold water. The cloth is run in this bath for 2 hours. A wash in a bath made slightly alkaline by adding trisodium phosphate, followed by a thorough rinse completes the process. It is sometimes necessaary to add a small amount of Acid Violet, Color Index No. 698, to the last rinse to obtain the bluish white which is usually requested.

Potassium permanaganate also has a limited use in producing novelty effects on shoe plush. The shoe plush after a good scour is dyed brown by running in a bath containing 30 lb. of permanganate per 825 gal. of water at 120° F. for 11/2 to 2 hours. An addition of 5 to 10 lb. of potassium permanganate is usually necessary to obtain the desired depth of shade. Following the dyeing the cloth is rinsed at 160° F. with water made slightly alkaline by adding 1½ lb. of trisodium phosphate. Two warm rinses complete this part of the process. The novelty two-colored effect is obtained by using a brush tipping machine. The latter is essentially a one-color printing machine which uses a brush roller instead of an engraved roller. The pile is tipped with an acidulated solution of hydrogen peroxide. If nothing more is added to the tipping liquor a brown pile with a lustrous white tip is obtained. By adding certain basic and acid colors not affected by the peroxide beautiful blue, green and rose tips over a brown base are obtained.

A gray, varying in intensity from a light rabbit's fur color to a jet black, can be substituted for the brown at the base of the pile. The depth of the gray is directly proportional to the depth of the manganese brown originally on the fiber. It is accomplished by immersing the cloth after the tipping treatment in a cold bath containing 5 to 12.5% andine salt and .25 to 12.5% sulphuric acid, depending on the depth of shade desired. It is worked in this bath for 30 minutes. A weak ammonia rinse and a thorough wash completes this process.

The above principle—aniline black over manganese brown—is sometimes utilized to obtain clear white discharges on woolen fabrics.

Bleaching Yarns, Skins and Straw U. S. Patent 1,966,915

One hundred grams of woolen yara may be placed in a solution of 1000 cc. of methyl alcohol in which 30 cc. of hydrogen peroxide (30% water solution), are incorporated. As the oxygen of the hydrogen peroxide is liberated much more freely in an alkaline solution, there should also be added about 2 cc. of, preferably, concentrated ammonia water. The solution containing the yarn should be heated to about 60° C. for about 8 hours.

The pelt is put into a bleaching liquor of about 1000 cc. of ethyl alcohol containing about 15 cc. of hydrogen peroxide (30% water solution), about 0.3 cc. of concentrated ammonia water, and about 45 g. of Turkey red oil. The skin is allowed to remain in the bleaching liquor for 24 hours at about 18° C. The skin thus treated exhibits perfect bleaching and the complete absence of injuries or impairments.

Pandan "stumps" are treated with a 1000 cc. ethyl alcohol solution containing 35 cc. of hydrogen peroxide (30% water solution) for about 6 hours, at about 60°C, and are then finished in the usual way. The bleaching proceeds very smoothly because the chlorophyl is extracted by the alcohol.

Natural Finish for Calico Potato Starch 5 lb. Wheat Flour 7½ lb. are boiled with Water 250 lb. then add

China Clay Paste 10 lb.

and

French Mineral White Boil and add	10	lb.
Coconut Oil	34	lb.
White Soap		lb,
Carbonate of Soda		lb.
Water	3	Bi.
Add to a vat containing		
Potato Starch	15	lb.
md		
Water	75	Њ.
Stir thoroughly and then	slowly a	dd
Potato Starch	5	llı.
nd		
Water	5	lb.

with a trace of ultramarine.

The statched goods are dried in a dry room, damped and rolled under pressure.

Alizarine Lake Formula No. 1

Sulphate of Alumina

(Tech. 18% Al₂O₃) 972 lb, Water 10,000 lb.

No. 2

Soda Ash 500 lb. Water 5,000 lb.

Filter both solutions.

Add the hot soda solution slowly to the hot alumina solution while stirring, keep boiling gently until the precipitate begins to be glassy, wash with clean water free from from until, by repentedly decembing, a sample of the wash water shows but very little turbidness with chloride of barium solution. The alumins now obtained by filtering may be used at once for making alizarine lake. The weight of the paste filtered into the bng amounts to about 7000 parts. Add to the alumina paste a solution of 144 parts calcium chloride anhydrous, chemically pure, in 500 parts water, and follow, while stirring well, with a solution of 84 parts phosphate of ammonia (pure neutral sult) in 500 parts water. Then stir in 150 parts ammonia Turkey red oil, which has been previously dissolved in a little water, and finally add 1000 parts Alizarine Red 1B extra (20% paste).

Either boil this preparation for 6-10 hours in an open vessel, when the evaporated water must be replenished, or treat for 1 hour in the autoclave with about 59 lb, pressure.

Alizarine Cyclamine is affected by metals including copper, and for this reason should not be steamed in the autoclave; lead vessels, however, may be used without risk.

Every substance used in the making of

madder lakes, including the water, must be free from iron.

Alizarine Dyeing of Silk

a. The well cleaned silk is entered, worked and steeped over night in a cold bath of basic aluminum sulphate prepared by dissolving 171/2 oz. aluminum sulphate, free from iron, in 1 gal. of solved in a pint of water is added, the clear solution showing 12-15° Tw. silk is wrung out from the mordanting bath, rinsed well, then fixed for half an hour in a cold bath of sodium silicate of 1° Tw. and finally rinsed very thoroughly. A basic aluminum salt is thus obtained on the fiber without injuring any of the properties of silk. The mordanted silk is then dyed with alizarine paste, the quantity of alizarine used depending upon the depth of the shade to be dyed, in a boiled off liquor bath broken with acetic acid, entering and working it in the cold for half an hour, gradually raising it to the boil in 1 hour and dyeing at that temperature for another half an hour. The dyed silk is then thoroughly washed in water, brightened in a weak bath of acetic acid and finally dried. The silk is dyed bright red.

b. Silk after being properly cleaned is entered, worked and steeped overnight in a cold bath of "nitrate of iron" basic ferric sulphate-32° Tw. It is wrung out the next morning from the mordant bath, rinsed well in water, then fixed by working for half an hour in a cold bath of sodium silicate of 1° Tw. and finally riused very thoroughly in water. This mordanted silk is dyed with alizarine as usual. This gives a bright violet color.

c. As chrome cannot be used with advantage on silk as with wool, on account of its tendency to destroy the luster and injure the fiber, the mordanting is usually done with chromium chloride or chromium sulphate. The well scoured silk is worked and steeped overnight in a cold bath of basic chromium chloride 32° Tw. The next day the excess liquor is squeezed out, the mordanted silk is well washed in water, fixed for half an hour in a cold bath of sodium silicate 1° Tw. and finally rinsed very thoroughly. A basic chromium salt is thus obtained as a mordant on the fiber without particular injury to any of the properties of silk. The mor-danted silk is then dyed in a boiled off liquor bath broken with acetic acid as usual. The dyed silk is thoroughly | tion), dye another 1/2 hour at 50-55

1

washed, brightened with acetic acid and dried. Silk is dyed a bright chocolate color.

Chrome Dyeing Formula No. 1

Chromium Black kg. Acetic Acid 4 kg.

Heat to 100° C., boil 1/2 hour, add: Formic Acid kg.

boil another 1/2 hour, add

Potassium Bichromate 1.5 kg. at a temperature of 70° C., then go up to 100° C., and boil 1/2 hour.

No. 2

Chromium Blue 4 kg. Acetic Acid kg. Glauber's Salt 10 kg.

Heat to 100° C., boil 1/2 hour, add Formic Acid

boil 1/2 hour more, add

Potassium Bichromate 1.5 kg. at 70° C. up to 100° C., boil 1/2 hour.

No. 3

Chrome Flavin kg. Additions, method as in No. 2.

Vat Dyeing

Formula No. 1

Vat Scarlet 4 kg. HN-Process, 50-55° C. "One bath" process. Fundamental Vat Dvestuff 1 kg.

Water 30 kg. Caustic Soda (40° B6.) 1 kg. 1 kg. Hydrosulphite

Dyeing Vat Glue 3 kg. Ammonia 3 kg.

Hydrosulphite Dye 1/2 hour at 50-55° C.

No. 2

2 kg.

6 kg.

Vat Scarlet HN-Process, 50-55° C. "Two bath" process.

Fundamental Vat As in No. 1.

Dyeing Vat
As in No. 1.

Dye 1/2 hour at 50-55° C., then add 2% hydrosulphite for the second (applica-

	TEXTIL
No. 3 Vat Black	2 kg.
HN.Process at 50-55° C. "One bath" process.	g.
Dissolve the solid vat (kur	pe fest) in
the same amount of boiling was 5% (of the dyestuff weight)	ter, adding
the same of hydrosulphite.	giue and
Dyeing Vat Gluo	3 kg.
Ammonia.	3 kg.
Hydrosulphite	2 kg.
Dye 1/2 hour at 50-55° C.	
No. 4	
Vat Black	12 kg.
HN-Process, 50-55° C. "Two bath" process.	
Solid vat solution (see No.	3).
Dyeing Vat	٠,٠
Glue	3 kg.
Ammonium Sulphate	4 kg.
Hydrosulphite	2 kg.
Dye 1/2 hour, then add:	
Hydrosulphite	2 kg.
Ammonium Sulphate	3 kg.
Repeat dyeing 1/2 hour (sec	oud bath).
No. 5	2 1.0
Vat Blue HW-Process, 60-65° C.	3 kg.
"One bath" process.	
Fundamental Vat (Stammküpe	٠)
Caustic Soda (40° Bé.)	2.2 kg.
Dyestuff	1 kg.
Water	30 kg. 1 kg.
Hydrosulphite	1 kg.
Dyeing Vat Glue	3 kg.
Ammonia.	3 kg.
Hydrosulphite	2 kg.
Dye 1/2 hour.	
Dyeing Formula for Acetat Velvet	e Rayon
Formula No. 1	
Substantive Dvestuff	2 lb.
Glycerin, Dynamite Condensed Water	4 lb.
Condensed Water British Gum Thickening	2 gal. 6 gal.
Caustic Soda, 75° Tw.	1 gal.
The following is an example	
color for acetate rayon velvet.	
Basic Color	1 lb.
Acetic Acid (30%)	20 lb.
British Gum or Senegal Thickening	8 gal.
A proportion of tannic aceti	
improves the fastness to washi	ng, in deep

shades.

The following is an example of a formula for a print color containing tanna act of the property of the color

Basic Color	2	lb.
Acetic Acid, 30%	15	lb.
Acetine	11/4	lb.
Water	5	gal.
British Gum	18	lb.
Tannic-Acetic Acid 1:1	10	lb.

The last named ingredient should be added only when the color has become cold.

After steaming, the pieces are treated for a few minutes in a lukewarm bath charged with 12 oz. of tartar emetic per 10 gal. of water. This operation is commonly performed in a star machine, but it may also be carried out in a winch apparatus where the more robust velvets are being handled. After being treated with tartar emetic, the batch is given a light rinse in cold water, after which the pieces are hydro-extracted.

Vat Printing Color

Paste Vat Color	10	lb.
Glycerin, Dynamite	31/2	lb.
Carbonate of Potash	14	lb.
Sodium Formaldehyde Sul-		
phoxylate	7	lb.
British Gum Thickening	7	gal.

The following recipe for a color for the brush printing of viscose rayon plush will furnish an indication of the proportions of substantive dyestuff and other ingrodients used in preparing the print colors:

Diphenyl Brown BBN Extra	8	oz.
Direct Orange G	3	OZ.
Chrysophenine G	8	OZ.
British Gum (Dry)	8	oz.
Glycerin	10	oz.
Phosphate of Soda	12	07.
Condensed Water	1	gal.

Wool Dveing

Indigo (20% Paste) Water	10 2.4	lb. gal.
Sodium Hydrosulphite (Powder)	2.5	lb.
Canatia Soda (78° Tw)	в	nt

The indigo and the water are intermingled first. To this mixture, the sodium hydrosulphite is added, gradually and with unceasing stirring. Finally, the caustic soda is introduced. The mixture is to be frequently stirred and its temperature maintained at 60° C. In about two hours, complete reduction may be expected.

T-1:-- (COC()

Indigo Fermentation Vat Formula No. 1

Indigo (60%)	20-	40 lb.
Woad	560-1	120 lb.
Bran or Sharps	30	40 lb.
Madder	10-	15 lb.
Lime	12-	25 lb.
Water	3	240 gal.
No.		
Water	21	60 gal.
Woad		5 cwts.
Natural Indigo (P	aste) 20-	40 lb.
Bran	•	5 buckets
Madder		6 lb.
Lime (In Slaked F	orm)	3 gal.
Lime (In Slaked Fo	rm) as	directed

The water is run into the vat and raised to the temperature of 135° F. The woad is now added and the liquor stirred several times till "pasted." The 3 gal. of slaked lime are stirred in and the whole left over night.

A representative British hydrosulphite vat for wool may be made up in accordance with the following tabulation:

	diation.
Water	1080 gal.
Ammonia (25%)	3.6 pt.
Hydrosulphite Powder	2 lb.
	2.4 gal.
Indigo Solution (20%)	2.4 gal.
Olue Solution (1:10) Indigo Solution (20%)	2.4 gal.

The water is run into the vat and the temperature brought up to 120° F. The indigo solution is stirred in.

At the beginning of each dyeing operation, add animonia, hydrosulphite powder and indigo solution. At the end of the day's run, add a little glue solution and 1.2 qt. of caustic soda (at 70° Tw.).

Printing of Animal Fibers U. S. Patent 1,962,601

Colored patterns fast to washing, light, perspiration, etc., are obtained by printing prechlorinated wool and silk with a thickened paste containing Indigosol, Leucosol, or similar water solvent derivatives of vat dyes and sodium nitrite, then steaming with wet steam at 99-100° C, for 7 minutes, and passing the fabric in open width through dilute sulphuric acid (50 g. [density 1.53] per l.) at 95°, followed by washing and oxidation with a solution at 35-40° containing (per l.) 1.5 g. of sodium persulphate and 2 g. of sulphuric acid (density 1.015) for 20 reintees

Dyeing Aged Black on Piece Goods The following is suggested: 120 lb. aniline salt, 10 lb. aniline oil, 35 lb.

sodium chlorate, 1/2 lb. copper sulphate, per 100 gal. liquor.

The goods are impregnated with this solution, aged and chromed.

The following is another method of dyeing an "ungreenable" aged black.

Two solutions are prepared:

a. 55 gal. of water, 45 lb. aniline salt, 131/2 lb. toluidine, 7 lb. acetic acid, 181/2 lb. sodium chlorate.

b. 181/2 lb. nitrate of iron, 76.6° Tw., 6 gal. water, 27 lb. of a solution of copper sulphate (2:10).

Mix 8 gal. of a with 1 gal. of b, and pad with this mixture. Age and develop as usual.

The following process is also recommended: The pieces are padded with the following solutions, which are prepared separately, mixed when cold, and made up with water to 100 gal. The padding liquor should stand at 12° Tw.; 120 lb. aniline salt are dissolved in 26 gal., 3½ pt. water; 5½ lb. copper sulphate are dissolved in 10 gal. water; 37 lb. 9½ oz. sodium chlorate are dissolved in 7 gal. 3½ pt. water; to this are added 4 gal., 6½ pt. aluminum acetate, 15° Tw.

The cloth should be impregnated in such a manner that it retains about its own weight of padding liquor.

After impregnation, the cloth should be dried as rapidly as possible at a low temperature, after which it is aged for 1 to 2 hours at a temperature of 92° to 96° F.

The aging is followed by chroming and soaping.

Cotton Printing Paste

Victoria Blue B 6 oz. Methyl Violet 4B 42 oz. dissolve in

Acetic Acid, 40% 1/2 gal.
Starch Thickening (1 lb.
Wheat Starch/1 gal.) 5 gal.

when cold add
Tannic Acid (4 lb./lgal.) 4/2 gal.

Crimping Cotton

Beautiful effects may be obtained by printing on a Gum Resist and subsequently passing the cloth through strong caustic soda. The dry content of the gum used as a resist is very important. A very highly converted British Gum is usually used and the strength will run 3-4 lb, per gal.

The greater the dry content of a gum

resist, the more effective is its power to resist the caustic soda. The latter will vary in strength from 25 to 30% according to the length of time the cloth is let he after immersing and squeezing and prior to washing out. For best results it as advisable to select a light weight cotton cloth and print a design that is largely composed of lines running parallel to the selvage of the cloth. The reason for this is that the shrinkage, for the most part, is that the similar, to the most period takes place in the warp. After printing, run the cloth through the strong caustic soda in a pad box and let set 1 to 2 minutes. Finally inse well with cold and hot water, hydro-extract and dry in a crepe diver. In dyeing grounds for this type of work it is best to select colors that will not be affected by the caustic seda. If crepe dyeing is possible then beautiful two-toned enects may be obtained by dyeing the cloth after crimp-

In dyeing the latter, the dyestuff will have much more affinity for that part of the cloth that has been attacked by the caustic and as a result this portion will come out much heavier. Other effects may be obtained by selecting printing colors that will develop in a steaming operation and that will work well with the Gum Resist. These colors are printed on with the Gum Resist and then steamed, padded with the caustic and finished as mentioned above. The final result is a crimp in the colored or printed portion of the cloth. By selecting dyed grounds that may be discharged it is possible to obtain a crinkle in the white portion of the cloth. A discharge is made up with the Gum Resist and upon printing and steaming the color is discharged at the printed part. After running through the caustic soda and finishing as mentioned above, it will be noted that the cump is in the white portion of the cloth whereas the colored portion is uncrimped.

Lacquer Printing of Cloth with Metallic and Pigment Colors

This type of work is largely being carried out on silk, rayon and celanese where excessive handling is at be avoided. The advantage of this type of princing is in the fact that finished goods may be printed, dried and shipped without any intermediate process of steaming, washing, etc. The colors are really in a sense painted on the cloth and the secret of the success of this type of printing hes chiefly in the softiness of the resultant print. Formerly bronze and pigment prints were extremely hars when printed

by this method but today the lacquers used have been highly developed and the prints are much softer in feel. Both cellulose acetate and nitrocellulose lacquers are used and the difference between the two is very slight as far as the resultant print is concerned.

Bronze or metallic prints are nowheres near as fast as the pigment class of colors. They tend to go dull on standing and will wash out in time. Pigment colors are extremely fast and will even stand a good subbing. In order to do a perfect job, the engraver, printer and colorist must work together. The engraving is very important as too shallow a depth will make the color stick in. The colorist must have the proper amounts of solvents in his printing paste, so that the paste will not div too fast in the engraving. The printer must run at a uniform speed so that the paste as worked . out by the colorist will give even results. Too fast a drying paste will make the color stick in, whereas too slow a de paste will not dry fast enough over dry cans. A nitrocellulose lacquer can be made by dissolving the dry nitrocellulose in a mixture of acetone and ethyl or methyl acetate. A cellulose acetate lacquer can be made by dissolving the dry substance in a mixture of alcohol, phenol and solvent amplithm. In using pigment pastes it is advisable to have them extremely finely ground in some solvent, such as acetone together with olive or castor. Proper grinding requires special equipment and this treatment is very essential for the best results.

Anthre Black Printing Paste
Yellow Prussinte of Potash
Chlorine of Soda Crystals
dissolved in
Hot Water
and added to
Anthre Balt
Previously dissolved in
Hot Water
1 pt.
and stired into
Starch Tragacanth Thickening
Printing Paste
5 pt.

Silk Printing Pastes Formula No. 1

Five pounds of good white starch and 5 lb. of white dextrin are smixed with 1 gal. of water, 7½ lb. of accete acid of 12° Tw., 2 lb. of olive oil and 2½ gal. of water are then added, and the whole

boiled into a paste. This will suit almost all colors.

No. 2

Five pounds of good white starch are mixed with 1 gal. of water, and 21/2 lb. of pale glue previously dissolved in 21/2 gal, of water are added, and the whole boiled up to a paste. After allowing to cool, add 5 lb. of acetic acid 7° Tw., and 2 lb. of olive oil.

Textile Printing Pastes

Formula No. 1

Wheat Starch Thickening Wheat Starch 12

Water

Boil and add

61/2 oz. Chlorate of Potash an dissolved add 付:数少百

Yellow Prossists of Potask

niline Salt oz. 1/2 tumbler Printing Pasto 1 gal.

No. 2

Copper Sulphate Black for Block Prints (Thickening)

Oz.

Chlorate of Soda 07. Copper Sulphate 21/2 oz. Wheat Starch 5 lb. Water gal.

.Boil together and when thickened add Gum Tragacanth I agale Boil further until an even tempere is pro-Printing Paste

Use 7 parts of thickening to 1 of aniline hydroclaride.

Silk Printing Color Resist

650 lb, 50 lb. 30 lb. lb.

Turpentine Oil

are heated together until they form a

thoroughly liquid mass.

This resist is printed on lukewarm either in the printing machine with very deeply engraved rollers, or by hand printing. For the latter purpose the above mass must be kept a little thinner by the addition of a little more turpentine oil.

After printing, the goods are sprinkled with fuller's earth, and then hung up for a few days at the ordinary temperature. When the resist is dry, the goods are washed in cold water and dyed in a told

Fancy Textile Printing "Resists" The following is a good and simple formula generally used by textile printers. It washes with water.

Formula No. 1

top Black or

1/4 OZ.

h to make into a paste he above thoroughly and

Tincture of Green Soap mix and add:

Concentrated Los Solution 5 drops Stir vigorously and keep in well corked

No. 2 Japan Color ⅓ oz. Raw Linseed Oil 1 oz. Boiled Linseed Oil 1/2 oz.

Washes off with oil solvents.

Powdered Castile Soap 1 oz. Hot Water 2 oz. dissolve thoroughly and add show card color.

White R phur Dyes Water

Heat until the gum and salf solved.

If the resist white is found to run when printed with heavily engraved rollers, it may be improved by the addition of 75 to 100 g. China clay per kg. of color: as a rule, however, this addition is not necessary with ordinary patterns.

Cotton Yarn Dye Resist

first impregnated with nic acid overnight, rning, and then immersed in a bar of stannic chloride takes on the property of resisting a dyes. Use about 3 lb. of tannic act 100 lb. of cotton yarn), and fix in a bar containing 2½ lb. of stannic chi id crystals. This tannic acid bath is un hot at the time the cotton is entered, but is cold by morning. The tin bath is used

Wax Resist for Woolen Yarn

Rosin	60 lb.
Yellow Beeswax	5 lb.
Mutton Suet	2 lb,
Spermaceti	3 lb,
Paraffin Wax	2 lb.
Turpentine	4 lb.

The above are heated together and the resulting paste is printed on the goods. Strew with fuller's earth to prevent sizing, and when dry, wet out the printer cold water and dye in the warm bath with the recommendation.

Acid and Alkaline Resists atment for Wool

U. S. Patent 1,964,934

Sulphite Cellulose Waste
Laquor (Lime Karley)
Magnesium Chloride

10 oz.

Stripping Sulphur Colors from Mixed Fabrics

A simple and yet very effective way of stripping sulphur colors on cotton in the presence of wool or worsted is as follows:

Prepare a cold bath of ½° Tw. chloide of hme. Run the cloth full width in his bath for 30 minutes. Then drop the ath and rinse thoroughly with cold atter. A second bath contaming ½° Tw. comparing hydrothoric yacid is now rade. The loth is run in this bath for manutes at 160° F, and then rinsed action of the companion of the results of the contamination of the cont

The chemic treatment should destroy tractically all the sulphur dyestuff inset of 15 minutes if the chemic is freshly nade. When old chemic solutions are seed, longer running or a stronger bath a necessary. The hydrochloric acid reatment removes any residue of rust or ulphide spots left from the chemic treatment. The resulting cloth is usually aght cream color which the sound of the considerably lighter. To the indicate the hidrinated slightly by the market considerably lighter. To the is hidrinated slightly by the market considerably lighter.

rosulphite Discharge on Indigo Ground
The printing paste is prepared as fol-

and printing paste is prepared as fol-

Hydrosulphite NF Concentrated 125-200 lb. are stirred into Hot Butish Gum

Thickening

655-580 lb.

and after cooling

Zinc White Paste 1:1 150 lb. Anthraquinone Paste 30% 50 lb.

Acetine (Neutralized with Soda)

are added.

20 lb,

The amount hydrosulphite in the discharge depends upon the depth of the indigo shade.

After printing, the goods are well dried at them stemmed for 3 minutes at 216-18 p.F. in the Mather-Platt, which must be referred to the matter of the washing anothine in a boding take containing 10 parts silicate of soda 66° Tw. to 1000 parts water and 3 performable by the washing machine in a boding the washing machine should take three-formths to one and a half minutes and the goods then well rinsed.

Instead of washing with silicate of soda, quick line (5 parts per 1000) or canstic soda solution may be used, although the subcate has the least effect on the indice bottom.

It is addisable to steam and finish the printed goods as quickly as possible, but if this cannot be done immediately, the naterial must be protected not only before but also after steaming against most air ke, winding rolls and keeping in a warm dry room 85-100° F. After steaming the white is cleared as above by passing the pieces this but alkalim.

Although the indigo is readily converted into a leuer compound by hydrosulphite, still the discharged places are up to show a bluish tint if the reduced compound is not completely removed from the printed parts, or if the indigo white is partly reoxidized, the fore the steamed process of the different parts of the indigo white is partly recoxidized, the continuous process of the pastern of updated in the discharge of hydrosultand reverse in indigo white from reoxidized.

Crease Proof Fabric British Patent 424,535

Ammonium sulphocyanide in the presence of variable quantities of urea has the advantage of requiring a comparatively low temperature for its formation. In previous similar processes it has been found necessary to heat the resin mixture for several minutes at 160 900 lb.

300 lb.

to 180° C. in order to produce full polymerization, but with these new resins a treatment of one minute only at 120° C. is sufficient; the textile material being treated is thus less liable to impoverishment.

The following is an example of the manner in which viscose rayon fabric is given a good feel and made uncrushable: First a solution is prepared, with the following ingredients:

30% Formaldehyde Urea 30% Ammonium Sulphocynnide Solution

cyunide Solution 150 lb. Wuter 900 lb.

The fabric is impregnated with this liquor, squeezed free from excess, and then dried. Afterwards the fabric is led over rollers heated to about 130° C. and the impregnated substances then react to form an elastic insoluble resin which makes the viscose fibers practically uncrustable.

It is possible to use an ammonium sulphide instead of the more expensive sulphocyanide and also to color the fabric during impregnation with the resin components. Thus viscose tayon fabric is impregnated with the following liquor:

30% Formaldehydo	900 lb.
Urea	300 lb.
30% Ammonium Sulphide	150 lb.
Sulphonated Cetyl Alcohol	
(Wetting and Dispersing	
Agent)	60 lb,
Ammonium Sulphocyanide	50 lb.
Diamine Sky Blue FF	20 lb.
and then dried at 150° C. for	10 minutes.

Crease sisting Fabric U. S. Patent 1,980,676

Fifteen gallons of casein solution containing 1 lb. of dry casein and 2 oz. of trisodium phosphate are mixed with 5 gal. of 30% later solution containing 2% zinc oxide on the dry rubber and 2% piperidine pents metalline dithiocarbamate. The latter material acts as an accelerator for the rubber. An ordinary sizing mangle can be used, the excess size being removed and the fabric is then dried. Subsequently, the fabric is washed in boiling soap solution to remove that part of the size which held the latex in suspension, presumably the casein component. In order to prevent the crossed yarns from adhering to one another, work the fabric during the drying operation which is the method employed in the acid organdic process for the same purpose.

Delustering Finish for Rayon

Fuller's Earth
 Titanium Dioxide
 40 lb.

3. Sulphonated Castor

Oil (30%) 150 lb. 4. Stearic Tallow Softener 15 lb.

Mix 1 and 2 and wet out with 3. Then add 4 and grind well.

Deguming and Decolorizing for Straw British Patent 424,189

Soda Ashar Rosin Casein 80 lb, 80 lb, 250–300 lb,

Water wive consistency of soft soap while being heated.

Renovating Surfaces of Textiles British Patent 419,856

The shine produced on textile fabrics by wear can be removed if the fabrics are first dry-cleaned, the surface fibers raised by teazelling, and then a mixture of 1 part sodium salicylate, 2 parts borax, 1 part cresol saponatis, and 3 parts ammonia in 320 parts water applied finally the goods are brushed thoroughly.

Mercerizing Wetting Out Agent U. S. Patent 2,008,458

Cresol, Technical Aniline 90 lb.

Mercerizing German Patent 606,025

As wetting agents for use in mercerizing lyes, use is made of acid esters of phosphoric acid in association with phenols and (or) highly sulphonated oils. A typical wetting agent comprises dibutyl ephosphate 1, crude cresol 9 and a highly sulphonated oil 2 parts by weight.

Low Luster Artificial Silk

Casein 10 lb.
Water 200 lb.
Turpentine 10 lb.
Petrolatum

10% of weight of cellulose

The above is emulsified and added to
the spinning solution (viscose).

Partially Saponifying "Celanese" To dye directly and uniformly with certain dyes, it is often necessary to partially saponify "Celanese" by padding with the following and drying.

Soda Ash 30 lb.
Glycerin 2 gal.

After drying, steam for 4 minutes in a rapid ager. Rinse well and dye with any direct dyestuff.

Restoring Luster to "Celanese" Pad with 28% acetic acid, tenter and dry under tension. Rinse well and dry.

> Rejuvenating Cloth U. S. Patent 2,006,192

A composition suitable for treating worn shiny wood or silk fabries is formed of alcohol 16 oz., 24% ammonia solution 3 oz., glacial acetic acid 4 oz., oil of lavender 1.5 g. and chloroform 2 oz.

Cotton Softener

a. Tallow
Caustic Potash (50° Bé.)
b. Water
When a is saponified, add b with stirring and stir until solidification begins.

Pre-Shrinking Trentment of Cotton Fabrics

U. S. Patent 1,959,406

Cotton fabric is shrunk by immersion for 1-10 hours in an aqueous liquor at 65-100° containing 1-4 oz. of ammonium alum and 0.25-3 oz. of sodium bisulphate per 10-50 oz. of water, followed by hydro-extraction (without intermediate washing) and drying.

Tarnish-Proof Cloth U. S. Patent 1,933,302

The cloth after dyeing is dipped in a solution of a cadmium salt (0.5 lb. or gal.) e.g., cadmium acetate which absorbs hydrogen sulphide when used as a wrapping for copper and silver articles and thus protects them from atmospheric tarnishing.

"Cravenetting" Textiles

The process of waterproofing or cravenetting proper is not a simple one. Soaking the fabric in a strong solution of acetate of alumina for several hours, extracting and allowing to dry slowly, is about as effective as any simple process. The acetate of alumina may be prepared by dissolving 1 lb. of alum in 1 gal. of hot water. In another vessel containing

½ gal. of water dissolve 1½ lb. of sugar of lead (lead acctate). Mix the two solutions and allow the precipitate to settle. The clear liquid only is used in preparing the bath, using about 1 qt. of the solution to 1 gal. of water.

Proofing Against Moth and Fungi British Patent 413,445

Animal fibers such as wool, felt, fur, skins, feathers, silk and hair, are proofed against moth and fungi by treatment with a solution of chromium fluoride so that a definite quantity of chromium compound equivalent to 0.65% of chromium fluoride is retained by the material. After steeping or padding with the aqueous solution, excess is removed and the chromium compounds fixed on the fiber by drying at a temperature above 150° F. In British Patent 418,529, the process in the above specification is modified by adding antimony fluoride to the chromium fluoride bath.

Mould and Fungi Proofing of Textiles British Patent 413,648

About 5% barium borate is claimed as an impregnant.

Silk Wool for Knitting

Silk wool, suited for knitting, may be produced as follows: The woolen yarn is first treated for 15 to 30 minutes in a cold bath of 100 l. in which % l. of hydrochloric acid (at 32° Tw. = 1.160 sp. gr.) has been dissolved. The yarn is now to be well drained or else hydrocxtracted. A second cold bath is prepared by using the clear liquor from a solution of 1½ kg. of bleaching powder in 100 l. of water. The yarn is treated in this cold bath for perhaps 15 to 30 minutes. Afterwards the yarn is drained and then soured with hydrochloric acid for 30 or 45 minutes. Next, the work is to be rinsed and then turned for 15 to 30 minutes in a warm bath at a temperature of 75° C. (167° F.). This bath is to contain 600 g. Marseilles soap per 100 l. of water. The work is now removed and hydrocxtracted. Afterwards, it is given a second souring with hydrochloric acid. Finally, it is well washed.

Felt Hat Stiffener Carnauba Wax Emulsion (Bright Drying) Shellac (Ammonia Water Solution) «

90 lb.

Stiffening Material for Shoes French Patent 777,404

The material is made by impregnating cloth, paper or felt with a colloidal substance, a part of which is in the precipitated state and consequently easy to dissolve while the rest is not precipitated and therefore less easy to dissolve. Thus, flannet is impregnated with a colloidal solution containing cellulose mitrate 150 kg, alcohol 580, acetone 60, carbon tetrachloride 120 l. and then dipped in water for 15 minutes. A part only of the nitrate is precipitated and the material is air dried.

Rubber Latex as a Textile Finishing Agent

The use of rubber as rubber latex or in a dispersed form has found many applications of late in the textile industry. It is natural to assume that a substance possessing the characteristics of rubber, i.e., water repellency and its flexibility, and especially the fact that it may be applied to a textile in a liquid state like many other finishing compounds, should find development in the finishing of textiles.

The application of rubber latex in connection with textiles has been grouped as follows: For the production of artificial leather and non-skid rug underlaps; as a backing and strengthening agent for fabrics that otherwise would be too sleazy for rough usage; for double texture fabrics. Hauser has discussed the use of latex in combination with canvas for friction belts, as well as its use as a binding agent for applying flocked wool or cotton to a fabric base.

The utilization of rubber latex in the carpet industry has assumed a rôle of importance as carpetings impregnated with it form their own selvedges without unravelling, thus obviating the necessity of a binding. Carpetings of this type may be joined together by use of a latex adhesive without any evidence of a surface seam. If the proper latex is used for the backing of the carpet, the latex is waterproofed to such an extent that it may be scrubbed on a floor without the moisture coming through. Rubber latex has been an important factor in developing a new type of construction in carpets and pile fabrics. In this process, a hair batt is laid on a latex-coated base and the fabric subjected to a vulcanizing process. In this particular development the use of looms for the pro-

duction of the carpets has been don away with entirely.

It has been stated that it is obvious that the textile mill is not equipped to develop the various latex compounds required. A textile plant possessing the facilities of the average sizing and finishing equipment and laboratory will probably be in a position to develop rubber latex as a finishing agent.

Rubber Latex

The presence of rubber latex as a processing agent has been made possible because of developments in prolonging its stability. Crude rubber latex, when stabilized with ammonia immediately following tapping, will withstand reversion or coagulation for the interval of shipping time until it reaches its destination, where it is subjected to further stabilization with ammonia. The rubber latexes are white to grayish in color, and are found occasionally with a yellowish cast. Latex, when freshly collected from the tree, may contain as high as 50% rubber, but following stabilization the rubber content will drop usually to 40% and under. In a number of cases, before selling, it is concentrated by various methods, or is compounded for a particular need.

The concentrating of latex is carried out by various processes, which may be subdivided as follows: (1) by creaming promoted by centrifugal force much in the same manner as a cream separator; (2) by filtration through unglazed porcelain while the latex is kept in movement; (3) by evaporation after the latex is stabilized by a non-volatile stabilizer like soap or sodium alginate.

Water dispersions of rubber differ from latex in that the latter at no time in its processing has been reverted to the solid state, but has been kept liquid since its tapping from the tree. The water dispersion, on the other hand, is a stable dispersion of congulated, smoked rubber, plus various compounding ingredients, effected by mechanical means. These have been marketed by a number of the leading rubber companies already com-pounded, and they exhibit properties similar to rubber latex towards other chemicals. They are usually less expensive than latex and greater efficiency may be obtained by their use because of the greater rubber concentration of the majority of dispersions when compared to the ordinary 40% latexes. Water dispersions of rubber usually yield softer films, but one of their drawbacks lies in that many of these are not as lightly colored as rubber latex and consequently will not yield the latter's clear films.

Rubber Latex with Starch

Rubber latex may be incorporated with a starch sizing to add flexibility and water resistance when padded to a fabric. Crude latex in admixture with a starch sizing will not waterproof a fabric but it will enhance its water repellency. However, if a compounded rubber latex is used, waterproofedness will be produced.

Rubber latex in mixture with starch is used extensively today as an adhesive. The mixture is not an easy one to produce. This is due to the action of a starch paste, which, although it is itself a protective colloid, tends to congulate latex when it is added in a hot state. The latex should be first protected with a protective colloid such as glue, casein or gum trageanth. Bone glue has been found to be an excellent protective agent as well as one exhibiting properties akin to a starch.

One part of a better grade of bone glue is heated, while stirring in 8 parts of water, to 140° F. until all lumps have been dispersed, and a smooth thin paste results. The glue should not be heated to over 140° F. since a decomposition of the protein may result.

If the cooked glue is tested for acidity it will be found to be somewhat on the acid side. Any substance exhibiting an acid reaction should not be added to rubber latex as acidity will tend to cong-ulate it. Consequently the glue is made alkaline with 0.5% solution of caustic soda, and cooled to about 110° F. (Although precautions against the addition of caustic soda to latex have been advised, no deleterious effects from the addition of small amounts of it have as yet been noted.) The latex-four parts of latex to one part of glue by volume—is then further stabilized with a small amount of ammonia, and then poured slowly while stirring into the glue. Thus we now have the protected latex mixture.

The starch (maize cooked 1 lb. to 1 gal.—tapioca starch 8 oz. to 1 gal.) paste is cooled to 140° F. and an equal volume of water is added. This should be made alkaline with a small amount of ammonia; the protected latex mixture is added to it slowly and stirred until a uniform mixture results. If this size mixture is padded on to a cotton fabric, a firm, flexible finish will result. Thus in a like manner it may be thinned to yield the desired firmness.

In adding a protected latex solution to

a cooked starch, care should be taken that the size should not be too hot-not over 140° F., since there is a hability of coagulation of the latex. Once a latex reverts or congulates, there is little hope for its redispersion, since this may be carried out only with special equipment as that used for making water dispersions of rubber. However, there are certain indications of partial coagulation before a latex will completely revert. If, upon the addition of the protected latex to the starch, a sudden stiffening of the latter is noted, we have an indication that congulation is setting in. No further addition of latex should be made, but the starch should be further thinned with ummoniated water until it thins out evenly, and then the remainder of the latex is added slowly.

A size-latex mixture as prepared above will produce a water-repellent finish on a fubric but will not waterproof it. To produce a waterproof finish, a "curable" or vulcanizable latex must be used.

Compounding Rubber Latex

In order to compound crude latex for vulcanization, there are certain essential chemicals which should be present in the inixture at all times. These are sulphur, zinc oxide, and an accelerator. Sulphur, cluoride may be substituted for sulphur. Any other chemicals added are for the purpose of lending some desired property to the resultant rubber film.

Any substance added to latex must be water-soluble and completely miscible with it, in order to produce effective results. Sulphur and zine oxide in their dry state are not soluble in water and therefore cannot be incorporated into latex as such. Sulphur chloride is miscible with latex, but because of its cost and its irritating action on the skin should he disregarded. Thus the zinc oxide and sulphur must be placed in a water-soluble state before their addition to latex. This is done by placing them in a colloidal stute, and they are marketed as colloidal sulphur and zinc oxides and capable of being thinned to a great extent with water before they fall out of solution. On a dry basis, the concentration of dry sulphur in the colloidal material is about 45% by weight, while the zinc oxide runs about 54% by dry weight.

The purpose of the sulphur in the mixture is to produce greater flexibility and toughness in the rubber film. To hasten this effect, zinc oxide is added. It may be termed a very slow accelerator in the vulcanizing or "curing" action of the sulphur on the rubber. However, to hasten the reaction between the rubber and sulphur to a greater degree, a more rapid outside accelerator is invariably added as well. Water soluble accelerators are present on the market which will cause the rubber to vulcanize at a temperature of 140° F., and it has been noted that latexes compounded with these accelerators vulcanize at oven temperature. A simple starting recipe for a vulcanizable rubber mixture is:

 Latex (50%)
 1
 gal.

 Colloidal Sulphur (45%)
 1¼
 oz.

 Colloidal Zinc Oxide (54%)
 2
 oz.

 Accelerator
 ½
 oz.

In preparing this mixture, the colloidal sulplur and zinc oxide are first thinned separately with a portion of the latex before their addition to the major portion. The accelerator is first pasted with a little sulphonated castor oil, and then thoroughly dissolved in a small amount of water at 150° F. The solution is then strained through a cheese cloth into the partially compounded latex. The latter is then stirred thoroughly to produce a uniform mixture.

If an accelerator which must be emulsified before adding to latex is used, it should be emulsified with triethanolamine and oleic acid as follows:

Accelerator	100 lb.
Oleio	5 lb.
Triethanolamine	2 lb.
Water	80 lb.

The accelerator and oleic acid are thoroughly mixed and added slowly while stirring to the triethanolumine diluted with the water. The amount of this emulsion added to the latex should be based on the actual weight of the accelerator present in a specific volume. For liquid accelerators, the dispersing of these in water with ammoniacal casein is recommended. An agitator must be used in order to obtain a stable dispersion. In order to prevent rubber films from oxidizing too rapidly, compounds called anti-oxidants are often incorporated into the latex batch. For the majority of water soluble anti-oxidants used with latex, the amount used is about double the weight of accelerator in the formula. If this vulcanizable mixture is protected with glue in the same manner as the crude latex and then added to a size batch which is applied and dried into a fabric, a complete waterproof should reault.

Care should be taken in drying fabrics impregnated with a starch-crude latex sizing on a can dryer. A crude latex film

when subjected to heat has a tendency to become soft and sticky, thus tending to adhere to the dry cans. If the percentage of latex in the size batch is such that this occurs, the sticking may be overcome by powdering the cans with a small amount of talcum. With a tenter dryer, little difficulty should be encountered in this direction.

In coating fabrics with latex for adhesive purposes or for producing pro-tective films, it is desirable that greater amounts of latex should be carried to the material. This is accomplished by use of a thickening agent on the same principle as the use of a thickener in printing fabrics. A more concentrated latex may be used alone since it is naturally creamy and thick. A natural 40% latex, how-ever, must be thickened. Thickening agents include starch, water soluble resins and colloidal clays. Where a coating is desired which overlooks the brittleness produced by the starch, then the latter should be used. Colloidal clays should be used when the natural flexible rubber films are sought. Much of the firmness as produced with a starch may be overcome by the addition of a softener such as sulphonated castor oil. If an excess of the sulphonated castor oil is used, tackiness in the crude rubber film re-

Of the clays, a good grade of colloidal bentonite makes an excellent thickening agent. A concentration of 1 lb. to a gallon of water in admixture with 1 gal. of crude latex yields a viscosity which produces continuous films having good body. To produce the clay paste, the dry bentonite should be first pasted with a small amount of sulphonated castor oil thinned with a portion of the subsequent water to be used. The remaining water is then stirred in and the mixture allowed to soak overnight for the lumpy clay to expand. On the following day the paste is thoroughly mixed and then strained through cheese cloth before its addition to be used. The remaining water is then stirred in and the clay will tend to dust when it is found present in the rubber film.

If it is desired that the film should be colored, an organic dye in solution may be added, but the greatest fastness is obtained by use of water-soluble dispersed colloidal pigments which are present on the market.

Films produced from crude latex mixtures, as pointed out previously, will tend to grow tacky with heat. If this condition is undesirable, a compounded latex must be used.

Wetting Agents with Latex

Recently, a number of wetting agents have been marketed especially for use with latex. These are of use when a thorough impregnation of a heavily woven cotton fabric is necessary. A wetting agent showing an acid reaction when in solution should be avoided. The best method of accomplishing a thorough impregnation of a heavy cotton fabric is first to boil it out thoroughly in sofa and in a wetting agent, and after a thorough wash and nipping it should be run through a pad in open width containing the latex and wetting agent.

the latex and wetting agent.

If the material is but wetted in water and the wetting agent added to the latex bath, then the high speed of the pad should be diminished. Instead, the cloth in open width is run very slowly through the latex in order to insure a thorough soaking, and then through the nip.

Precautions in Handling Latex

- (1) There should be a word of advice to the workman handling latex, and thus is that he should abstain as far as possible from placing his hands in the raw latex. The reason for this is that in many cases there is an acidic reaction from the perspiration on the hands which tends to cause reversions. Cases of latex congulation have been reported due to this cause.
- (2) Rinds and latex films that are noted on the surface of a latex bath should be picked off, since these hasten coagulation. If possible, when these occur, the bath should be strained through a cheese cloth to remove the films.
- (3) Latex should not be subjected to abnormal conditions of temperature. Latex when frozen will coagulate when reliquefied, and consequently should never be stored in a spot where a low temperature of 32° F. may occur. Latex should not be heated as this will cause the stabilizing ammonia to volatilize, this condition tending to hasten coagulation.
- (4) Latex mixtures should not be made in copper vessels, since if small amounts of copper are present in a rubber film the metal will tend to hasten the oxidation of the film.
- (5) Latex should never be added to size baths containing calcium, barium, or aluminum salts, as these exert a coagulation action on latex.

Rubber latex has found a place for itself in the finishing of certain textiles. It can be handled properly with the finishing equipment of the average mill. The prime requisite is that the finisher familiarize himself with this somewhat new finishing agent.

Fireproofing Solutions

The following is the formula of a solution used in theatrical work for rendering materials non inflammable:

Tungstate of Sodium 1742 oz.
Water 145 pt.
Dissolve in the cold and add:
Sodium Phosphate 242 oz.
Water 1 pt.
or a sufficiency of water to make the so-

lution sp. g. 1.140.

Dip the material in the solution, wring out with the hands, dry, and iron if necessary.

The following are formulæ of solutions advised by the L.C.C. for rendering curtains, Christmas decorations, etc., non-inflummable:

Formula No. 1 Ammonium Phosphate lb. Ammonium Chloride 1b. 11/2 gal. Water No. 2 10 oz. Borax Borie Acid OZ. 1 Water gul.

Beth solutions can be used for coarse fabrics, but No. 2 is better for more deheate articles. The fabrics should be dried without rinsing, and it is advisable to experiment with a small portion of the cloth before treating the whole, as the texture and colors of some materials are affected detrimentally.

Fireproofing for Canvas

Ammonium Sulphate	8	OZ.
Ammonium Carbonate	2.5	oz.
Borie Acid	3	OZ.
Borax	2	OZ.
Starch	2	OZ.
Dextrin	0.4	
Water	100	oz.

Steep 1/2 hour at 86° F.; 2 dips necessary for best results.

Fireproofing Brake Lining U. S. Patent 2,001,194

Brake lining is impregnated with a composition such as may be formed from an aniline dye 10 to 20 g., ammonium sulphate 60 lb., ammonium phosphate 10 lb., boric acid crystals 12 lb., gum acacia 2 lb., cresley ore 2 lb., barium hydroxide 4 lb., aqueous ammonia 1 qt., ammonium alumium sulphate 2 lb., copper-sodium

5-6 gal.

alginate 1.5 lb., benzaldehyde 1 oz., sodium bicarbonate 2 lb. and water 100 gal.

Flameproofing and Fireproofing Textiles
Sodium Borophosphate Resin

(Abopon) Water

Dip the textile into the above solution warmed to 110 to 170° F; wring out and pass between warm rollers. This process gives a uniform coating which does not powder out like the usual fireproofing salts.

Waterproofing Canvas Formula No. 1

A treatment that is sometimes given to awnings to waterproof them and still leave them flexible so they can be rolled up and down, is as follows: First apply a coat of glue size, made by dissolving 1 lb. of high-grade glue in 3 qt. of water. To 1 gal. of this size add 1 oz. of alum, previously dissolved in hot water. Apply the size while still quite warm, using a wide flat wall brush. When the size is dry apply two coats of a paint made by mixing white lead-inwith necessary tinting colors added, thinned to rather stout brushing consist-ency with a liquid composed of 2 parts of boiled linseed oil and 1 part of turpentine. Be sure to use boiled linseed oil, as raw oil would have a greater tendency to rot the canvas, more especially if glue size has not been used under the paint. Two coats, or not more than 3 coats, should be sufficient. Be sure to allow ample time between coats for thorough drying. If the use of paint is objectionable, shave paraffin into gasoline, in the proportion of 2 oz. of paraffin to 1 gul. of gasoline, stirring until the wax is dissolved. The wax must be in very thin shavings to dissolve quickly in cold gasoline. As soon as the wax is dissolved, brush a coat of the solution on the bare canvas, using a wide flat wall brush. The next day another coat may be applied. If you brush the material on carefully you should be able to build up a reasonably smooth, waterproof surface in this way. Be very careful when using this preparation that no one strikes a match near you, and that there is no sort of flame in the room where you are using the so-One of these processes embodies the use of paint and the other a wax as the

waterproofing agent, and either will leave the canvas reasonably flexible and water proof.

No. 2 Canvas Waterproofing

Waterproofing Cotton Cloth

Pad the cloth with aluminum acetate solution (2° Tw.) and dry. Then immerse in sodium stearate "solution" (5%) at 120° F. Rinse well and dry.

Tarpaulin or Tent Waterproofing Formula No. 1

British Patent 414,242

Paraffin Wax 3-5 lb.

Naphtha 200 lb.

Warm together on steam bath and mix until clear. Then mix in:

Aluminum Powder 5-20 lb.

Australian Patent 17,598

Water-Repellent Fabric U. S. Patent 1.967.267

Fabric is impregnated with a solution of 1 pt. of wax (or animal and vegetable fats, greases, or oils) and 1 pt. of water shedding substance (e.g., cellulose acetate or nitrate, etc.) in an organic volatile solvent (e.g., ethyl nectate) and then dried, whereby it retains its original softness but becomes water repellent.

Textile Backing (Waterproof)

Latex (50% Concentration) 1	gal.				
Casein	12	oz.			
Water	1	qt.			
Zinc Oxide	1½	oz.			
Sulphur	5	oz.			
Accelerator No. 552	½	oz.			
Agerite White Powder (Anti-Oxidant)	5	oz.			
Casein	Casein	Casein	Casein	Casein	Casein
Casein	Casein	Casein	Casein	Casein	Casein

Waterproofing Wool Goods

The simplest method of waterproofing wool goods is the application of metallic

salts and tannic acid, sold either as powder or crystallized, with or without previous or subsequent soap, or fatty acid baths.

Formula No. 1

For 100 l. of impregnation bath there is dissolved about 100 g. of acetate of lead, 200 g. of alum, and 100 g. of tannin in boiling hot water. The goods are passed at about 40° C., centrifuged, and dried at from 40 to 50° C. The effect of the impregnation process is considerably increased by the above-mentioned soap and fatty acid baths.

No. 2

Three hundred grams of the best sulphonated oil, and 100 g. of olive oil soap are stirred in 10 l. of boiling water. They are added to a bath of 90 l. water at a temperature of 50° C. and the goods are passed at 40° C. To simplify the procedure these two baths may be combined in one.

No. 3

One hundred grams acetate of lead, 200 g. alum, 100 g. tannin, 20 g. hinseed oil, 500 g. Monopol oil, and 100 g. of pyridine are well stirred into about 20 l. of boiling water and brought to a boil again. Then the whole is increased to again. Then the whole is increased 100 l. by adding water of at least 60° C. The goods are passed at 40° C. and dried rapidly. Wool fat that can easily be emulsified is also well suited for the wet impregnating of wool. When it is used, the emulsifying is done separately.

No. 4

Ten kilograms of wool fat, 1 kg. ammonia, 5 kg. sulphonated oil, and 500 g.
pyridine are brought to the boil in about
50 l. of water, the whole being well stirred. This suffices for an impregnating bath of about 800 l. Into this bath, before adding the emulsifying agent, there are stirred 500 g. of pyridine, and the temperature is brought to 50° C. goods are dipped at from 30 to 40° C., centrifuged, and dried thoroughly. To make the impregnation more effective, there may be added to these baths tannin substances or metallic salts. effect is always superior when they are used in separate baths.

Waterproofing Wool, Silk, Rayon and Cotton

Examples for impregnating fabrics and wearing apparel of wool, silk, rayon and cotton are as follows: In 100 l. of petrol or other volatile hydrocarbon solvent, are dissolved by stirring well, 1

kg. of linseed oil varnish and 2 kg. of ceresin, the latter first being melted. The goods are thoroughly dipped, centrifuged, and dried in the open air. Subsequent steaming gives further assurance of even and thorough impregnation throughout the fabric. Fabrics can be steamed on a wet pressing roller. With very light colored and with white goods, the best wool fat is used instead of the linseed oil, and white paraffin instead of ceresin. Wool fat is recommended especially for wool goods when a soft feel is to be preserved, since after the admixture of varnish, the goods grow harder with time. The varnish impregnation is particularly suitable for coarser goods for which very thorough waterproofing is desired, especially for tentings, army blankets, weter pails, and for colored umbrella fabrics of all kinds of fibers.

Porous Cloth, Waterproofing

For this purpose a solution of acetate of alumina or acetate sulphate of alumina, which is prepared as follows, is chiefly used.

Sulphate of Alumina 665 lb. dissolved in

600 lb. Water 945 lb. Sugar of Lead dissolved in

Water

Dissolve each by itself hot, precipitate cold, draw the clear solution off and make to Twaddell 15°. In this manner a standard alumma sulphate acetate is obtained of which the greater part is deposited on the fiber in drying.

As woolen and half wool goods still contain some soap from the milling process, a soap passage is as a rule not necessary before impregnating with alumina; otherwise the goods are passed through a weak soap solution (3:1000), squeezed and dried without rinsing.

The goods are impregnated on a hank washing or open width washing machine provided with pressure rollers, by passing the dry goods for I hour through the diluted acetate sulphate of alumina of 3% Tw. (undried goods at 712-15° Tw.). The goods are then slightly centrifuged without rinsing or squeezed and then dried.

For wool and half wool goods a single impregnation will suffice in most cases; if a higher grade of waterproof finish is desired the treatment is repeated, inserting a soap passage if necessary.
In place of acetate sulphate of alu-

mina, formate of alumina may be used

with advantage. The latter possesses the advantage over the former that the danger of the subsequent tendering of the cotton warp in half wool goods, due to the formation of sulphuric acid in the fiber, is climinated. Formate of alumina is used in the same manner as acetate-sulphate of alumina.

Waterproofing and Fireproofing Fabrics, Paper, etc.

Austrian Patent 136,953

The material is coated or impregnated with an alcohol solution containing a reein, fat or like substance and a non-hydrolyzing salt of a metal of the 2nd periodic group which forms a colorless or transparent compound with the alcohol. A typical solution comprises resin 2, castor oil 0.5, crystalline zinc chloride 3, crystalline magnesium chloride 5, and 96% alcohol 12 parts. The solution may be applied to crepe paper.

Waterproofing and Flameproofing U. S. Patent 2,003,148

A method of compounding a composition of matter for flame and waterproofing aqueous cellulose media and their de rivatives comprises heating 640 part And the state of t ceases, adding 20 parts of boric acid previously dissolved in 128 parts of boiling water, adding 16 parts of borax and thoroughly mixing, adding 16 parts of starch previously cooked to about 1° Bé. and thoroughly mixing in the same under constant stirring; dissolving 6 parts of suitable soap in 128 parts of water and bringing it to the boiling point, thereafter adding the same to the previously compounded materials, bringing about emulsification of the whole and then lowering the temperature to 110° F. and digesting for about 2 hours thereby forming a first composition; bringing 640 parts of water to the boiling point and dissolving therein 80 parts of ammonium chloride, 48 parts of boric acid and 16 parts of borax in the order named, and each after the preceding has been completely dissolved, stirring has been completely dissolved, stirring the same thoroughly after all three have been added and dissolved, separately dis-solving 32 parts of soft gelatin in 250 parts of water and heating to about 200° F. under constant stirring and

thereafter optionally adding thereto 13% parts of glycerin, stirring thoroughly and then adding the same to the ammonium chloride-boric acid-borax solution under constant stirring for about 30 min. utes and then digesting for about 1 hour at about 140° K; dissolving 3 parts of suitable soap in 128 parts of water, heat-ing to boiling and adding 8 parts of dex-trin, stirring such constantly to insure uniformity and then adding such to the ammonium chloride boric acid borax gelatin solution, thereby forming a second composition; bringing 128 parts of water to the boiling point, dissolving therein 15 parts of soap bark and filtering, thereby forming a third composition; dissolving 32 parts of alum in 256 parts of water as a fourth composition; digesting each of the four compositions for about 4 hours while stirring from time to time; combining the first, second and fourth compositions in a common vessel and then adding the third composition under vigorous stirring.

Colloidal Textile Oil Formula No. 1

Castor Oil	20	gal.
Coconut Fatty Acids	100	gal.
Caustic Soda Solution		•
BEL	15	gal.
Water	30	gal.
Manipulation: Mix in the	order	given
C.		_
No.		
Castor Oil	15	gal.
Optorus Fact Acids	75	gal.
Water and	221/2	gal.
Calstic Soda Solution		
(30° Bé.)	111/2	gal.
Paraffin Oıl (28° Bé.)	82	gal.
Manipulation: Mix at 40°	J.	_

Colloidal Olive Oil

Continual Cityo Ci		
Commercial Olive Oil	90	lb
Caustic Potash Solution		
(32° Bé.)	13	lb
Water	150	lb
30 1 1 1 0 01 11		

Manipulation: Stir the caustic potash solution into the olive oil at room temperature and allow to stand overnight. In the morning add the water (which is previously brought to a boil). The mixture is well stirred during addition of the water, which is added slowly.

Acetate Rayon Oil Sulphonated Castor Oil (65%) 50 gal. Commercial Olive Oil 45 gal.

650 lb.

Acetic Acid (28%) 20 gal. 5 gal. 100 gal. Paraffin Oil (28° Bé.) Water

Manipulation: Mix the three oils and the water at 40° C. Then cool to 30° C. and stir acetic acid into mixture slowly.

Hosiery Oil

Sulphonated Castor Oil (65%) 1000 lb. 300 lb.

Caustic Soda Solution (27° Bé.) Water

Manipulation: Mix caustic soda solution with oil at 40° C., then add water slowly, maintaining temperature at 35-40° Č.

Kier Penetrant Oil

10 gal. Sulphonated Castor Oil (62% T.F.M.) 20 gal. 20 gal. Water

Manipulation: Sulphonate the castor oil to 62% T.F.M., settle and draw off. Mix in xylol first and then water, with agitation, at 35-40° C.

Silk Oil

Sulphonated Castor (58%)Paraffin Oil (28

Caustic Soda (27° Bé.)

Water Steam Distilled P

Manipulation of Sil dients in order named at 35-40° ing careful to add caustic soda solution and pine oil very slowly, with constant stirring and allowing mixture to cool to room temperature as the pine oil is being added.

Soluble Oil

Formula No. 1

Paraffin Oil (28° Bé.) Sulphonated Castor Oil 33 gal. (75%) 33 gal. Sulphonated Red Oil (75%) 33 gal. Manipulation: Mix at 40° C.

No. 2

Steam Distilled Pine Oil 50 gal. **Sulphonated Castor Oil** 50 gal. (75%)

Caustic Soda (27° Bé.) 10 gal.

sulphonated castor oil, then add the caustic soda gradually with agitation, maintaining the temperature noted above with constant agitation. When nearly clear solution is obtained add the water slowly, continuing agitation, then allow to cool rapidly.

Soluble Textile Oil

Xylol or Toluol 15 gal. Paraffin Oil (28° Bé.) 78 gal. Double Pressed Red Oil Alcohol 3 gal. Caustic Soda Solution (27° Bé.) gal. Water 1 gal.

Manipulation: Mix the paraffin oil and red oil, heat to 40° C., add the previously mixed water and caustic solution, then add the xylol slowly and the alcohol last and rapidly cooling them as quickly as possible after mixture is uniform.

Wool "Soluble" Oil

U. S. Patent 1,965,935

An oil such as a mineral oil 64, is used in admixture with "Carbitol" 2, corn oil soap 14, rosin 10, water 6 and diethyleno glycol 4%.

Neutral Light Mineral Oil

Double Pressed Red Oil No. 1 Lard Oil Manipulation: Mix at 45-50° C.

Equipment required: Wooden or lead lined mixing tank.

Paraffin Oil (28° B6.) 90 gal. 5 gal. Double Pressed Red Oil No. 1 Lard Oil 5 gal. Manipulation: Mix at 45-50°

Textile Sizing Oil

Sulphonated Castor Oil (62% T.F.M.) 800 1ъ. 550 lb. Water Caustic Soda Solution (27° Bé.) 350 lb. Silicate of Soda Solution (37° Bé.) 1300 lb.

Manipulation: Heat the sulphonated oil to 35-40° C. and slowly add the other Water 40 gal.

Manipulation: Heat the pine oil to 38° C. in the lead lined tank, add the lead li

Oiling for Viscose Yarn

Ammonium Oleate	100 g.
Oleic Acid	25-30 g.
Alcohol	15 g.
Apply at 40-60° C.	

A 1% solution of above works well at 40° C.; treating time 25 to 30 minutes.

Rayon Yarn Lubricant U. S. Patent 1,979,188

Mineral Oil	60	lb.
Triethanolamine Olcate	9.7	lb.
Mineral Oil Sulphonate	9	lb.
Potassium Oleate	16	lb.
"Carbitol"	5	lb.
Aniline	0.3	lb.

Synthetic Neat's Foot Oil

Extra Lard Oil	30	gal.
No. 1 Lard Oil		gal.
Light Mineral Oil	30	gal.
Manipulation: Mix at 40°	C.	

Rayon Identification (Revised Method)

The following systematic scheme, when carried out in the given sequence, serves for the rapid identification of rayons. This method can be depended upon by an experienced analyst, particularly when used in conjunction with filament count and microsopical characteristics. For the inexperienced man we recommend the detailed method of Rayon Analysis, and in comparison of the unknown rayon with standard samples of known make. The standards should be as inclusive of the rayon field as possible and should be kept up to date.

Rapid Method

Test 1-Identification of Animal Fibers

Millon's Test

Animal fibers—real silk, wool and hair—are quickly and positively identified by means of Millon's Reagent (see Identification of Rayon—Detailed Method).

Test 1A—Identification of Animal Fibers, Cellulose Fibers and Cellulose Acetate

Flame Test

Twist five or six strands of the unknown sample into a long, compact mass. Push the end of sample gently toward a match flame. (Do not allow sample to actually touch the flame.) Animal fibers tend to fuse and burn

Animal fibers tend to fuse and burn slowly when brought near to a flame. If the flame of the burning fibers is extin-

guished, the odor of the white fumes which subsequently arise from the smoldering end will have a "burned hair" odor. The burned ends of the fibers will have a dark, hard, brittle knob of material. Heavily mineral-weighted silks will leave a distinct ash which more or less retains the shape of the original material.

Vegetable fibers and most rayons do not fuse in the burning. They burn rapidly, and the fumes coming off after the extinguishing of the flame smell like burning cotton. Acctate rayons, in burning, smell like cotton and melt like animal fibers. The fused knob remaining after the flame is extinguished is hard but not brittle. If heated to a sufficient degree (in an evaporating dish or other suitable container) acctate fibers will melt without burning.

The burning test, while helpful, is not as instructive as the Millon's Reagent Test, inasmuch as it does not show the relative quantities and locations of the animal and vegetable fibers in mixed yarns or fubrics.

Test 2-Identification of Cellulose Acetate Rayon

Solvent Test

Cellulose Acetate Rayon is soluble in acetone; also in boiling 40% acetic acid. (See Identification of Rayon—Detailed Method.)

Test 3—Identification of Nitrocellulose Process Rayon

Diphenylamine Test

Nitrocellulose Rayon is turned blue by treatment with a solution consisting of 1% by weight of diphonylamine dissolved in concentrated sulphuric acetic acid mixture. (See Identification of Rayon—Detailed Method.)

Test 4-Identification of Viscose and Cuprammonium Rayons

Wright's Stain Test

Wright's Stain Test solution colors air-dried Cuprammonium Rayon violet and air-dried Viscose Process Rayon blue. (See Identification of Rayon—Detailed Method.)

Detailed Method

Chemical Identification of Rayon

(1) Identification of Animal Fibers in Mixed Fabrics

Millon Test

As small quantities of animal fibers present in unknown samples may cause

confusion in some of the following tests, an unknown sample should first be tested for the presence or absence of animal fibers. These are easily and quickly identified by means of the Millon Test, details of which follow:

Preparation of Millon's Reagent

Millon's Reagent is prepared by dissolving a given weight of metallic mercury in its own weight of pure concentrated nitric acid at room temperature in a non-corrodible container (porcelain, glass, agate, etc.). When completely dissolved, the solution is diluted and mixed with an equal volume of cold water. The solution should be clear.

If a yellow turbidity develops in the above noted solution, stir in a small quantity of nitric acid until the solution clears

Each new batch, or one which has stood open to the air for a long time, should be tested for proper activity by matching to the skin or by use of white animal fibers. When stored in air-tight glass stoppered bottles, the solution keeps for months.

Use of Millon's Reagent

Moisten the unknown swatch with Millon's Reagent. Warm to blood heat (do not boil) for a few seconds, or allow to stand for a few minutes at room temperature.

Animal fibers turn red, thus showing both their presence and position or dis-

tribution throughout the pattern.

Nearly all dyed animal fibers show an observable change toward red in this test without previous stripping of dye.

Swatches wet with water or with alcohol appear to react normally if flooded with reagent (to dissolve first precipitate).

Caustic Test

As minute quantities of cellulose and rayon fibers present in unknown samples largely composed of animal fibers may not be detected by the Millon Test, we recommend a subsequent caustic test for fabrics that appear by the Millon Test to be composed largely or entirely of animal fibers.

Although strength of solution, time and temperature may be varied over wide limits we recommend a 10% solution of caustic soda at 180° F. for 10 minutes.

Animal fibers dissolve completely. Cellulose and rayon remain in fiber form. (Note-Cellulose Acetate is partially saponified; regenerative cellulose fibers soften and dissolve to a limited extent. Cellulose Acctate fibers may be removed previously to caustic boil by use of ace-

(2) Identification of Cellulose Acetate Rayon

(a) Acetone Test

Place yarn or fabric in U.S.P. acetone

Cellulose Acetate is very readily dissolved.

So called "iron-proofed" Cellulose Acctate partially dissolves and leaves a leathery residue.

All other Rayons are unaffected by this treatment.

(b) Acetic Acid Test

Place yarn or fabric in a boiling solution of 40% acetic acid (C.P. acid is not necessary).

Cellulose Acetate is very readily dissolved.

So called "iron-proofed" Cellulose Acetate partially dissolves and leaves a leathery residue.

All other Rayons are unaffected by this treatment.

"Tron-Proofed" Cellulose Acetate
"Iron proofed" Cellulose Acetate is Cellulose Acetate that has been treated with an alkaline medium in such a way that the outside
of each individual filament is partially saponified

profiled "Actales yarm may be pressed "Iron-proofed" Actales yarm may be pressed "Iron-proofed" Actales yarm may be pressed actale, between the proof of provided actale insulates the unified or partially aspossible to actale insulates the unified actale insulates the unified proof of the pr lose Acetate.

(3) Identification of Nitrocellulose Rayon

Apply one drop of diphenylamine solution" to the dry unknown sample.

Nitrocellulose Rayon immediately turns a deep blue color after which it slowly dissolves to form a blue solution.

Other rayons are not colored blue.

All nitrated fibers-for example, Viscose Process Rayon nitrated for the production of special effects—show a blue reaction with diphenylamine solution. Many dyestuffs show a blue coloration when exposed to diphenylamine solution.

Nitrocellulose samples that have been stripped in a strong reducing bath will sometimes fail to give the blue coloration

Diphenylamine solutions is prepared as follows: Mix 66 g. concentrated suiphuric acid with 33 g of glacial acetic acid, then add 1 g. diphenylamine.

described above, however, their crosssections remain unaltered in shape.

The only positive test for Nitrocellulose Rayons is a microscopic examination.

(4) Identification of Viscose and Cuprammonium Rayon

(a) Wright Stain Test

Prepare a saturated solution of Wright Stain (dry powder) in denatured alcohol (95%). Immerse air-dried unknown sample into boiling Wright Stain solution and boil for a few seconds. Rinse the sample thoroughly in cold water.

Viscose Process Rayon is stained blue by this treatment.

Cuprammonium rayon is stained violet. (b) Schreiber-Hamm (Sulphide) Test

This test is suitable only for raw rayon of standard manufacture. Certain experimental yarns and processed yarns cannot be positively identified by this

A 5.g. sample of the unknown rayon (Viscose or Cuprammonium) is placed in a flask together with 100 cc. of water and 3 cc. concentrated sulphuric acid. The mouth of the flask is covered with a piece of lead acetate paper and allowed to stand on a moderately boiling steam bath for 4 hours.

If the sample is Viscose Process Rayon, the lead acetate paper will be stained brown or black.

If the sample is Cuprammonium Rayon, no discoloration should be observed.

(5) Identification of Undesulphurized Viscose Process Rayon

The difficulty of visually distinguishing between some delustered rayons and undesulphurized Viscose Rayon has sometimes led to confusion and improper rayon identification.

Undesulphurized Viscose Process Rayon can be readily identified by means of sodium plumbite solution.

Preparation of Sodium Plumbite Test Solution:

- (1) Dissolve 40 g. lead nitrate in 200 cc. of warm water.
- (2) Dissolve 70 g. of caustic soda in 300 cc. of water.
- (3) Add the caustic sods solution to the lead nitrate solution.
 - (4) Filter.
 - (5) Dilute to 2 1.

Method of Testing

A small quantity of the solution prepared as above is brought to the boil.

The unknown rayon sample is inserted into the boiling test solution for a period of 1/2 minute.

Undesulphurized viscose process varn turns black.

Incompletely Incompletely desulphurized viscose process yarns are turned black, dark brown, or medium brown, depending on the degree of desulphurization.

Desulphurized viscose process yarn is stained a brownish yellow color.

When possible, check tests on known samples should be run simultaneously with the test.

Microscopic Identification of Rayon

As rayons are most easily, quickly and positively identified by means of a microscopic examination, this method should be used whenever possible.

A microscopic examination of rayon is very simple and can be successfully carried out by men previously unfamiliar with the use of the microscope after a

few hours' practice,
For the benefit of those unfamiliar with the microscope and its use, we are pleased to describe the cheapest type of microscope that is, in our opinion, suitable for the microscopic examination of rayon. The analyst will need:

1. Microscope Stand and Lenses.

The instrument should be capable of magnifying to 400 diameters.

The above combination includes achromatic objectives, 16 mm. and 4 mm., eye piece 5× and 10×; and Abbe condenser N.A. 1.20.

- 2. Microscope Lamp 3. Microscope Slid Slides and Cover Glasses.
- 4. Mounting Medium (Methylene Iodide, or Monobromnaphthalene).
 - 5. A piece of thin glass rod. 6. A small scalpel or sharp knife.

Treatment of Viscose Products Austrian Patent 138,007

Rayon and other products made from viscose are bleached and desulphurized by treatment first with an alkaline solution of hydrogen peroxide at a low temperature and then with an alkaline solution not containing hydrogen peroxide at a raised temperature. Thus, rayon may be treated at atmospheric temperature be treated at atmospheric compensation with a solution containing hydrogen peroxide 0.5 and sodium pyrophosphate 1%, freed from excess of liquid, left to stand for 3 hours at 35° C, and then treated at 95° with a solution containing sodium pyrophosphate 1 and Marseilles soap 1%. Alternatively, the material may be

treated with a single alkaline hydrogen peroxide solution first at a low temperature and later at a raised temperature.

Preservation of Ropes

Make a solution of sulphate of copper (blue vitriol) in water, using 1 lb. of the crystals in 4 gal. of water and soak the ropes in this solution for 4 days, then dry them. The ropes will become impregnated with the copper sulphate, which will keep them from being attacked by parasites and prevent rot. The copper salt may be fixed in the ropes by the application of a soap solution, made by slicing 1 lb. of yellow laundry soap in thin slices and dissolving it in boiling water. Use 1 lb. of soap to a gallon of water. While the soap solution is still lukewarm put the ropes in it and let them soak overnight. Next morning take the ropes out and let them dry. The copper soap thus formed is more effective than tar, which is used on ropes employed by sailors, but tar is likely to stain painted surfaces, so painters should stick to the soap treatment. Ropes must be kept in a warm, dry place, never in a basement, because dampness would injure them in time.

Sash Cord Impregnants Formula No. 1

Paraffin Wax (130-

Caustic Soda

Ammonium Sulphate

132°F. M.P.)		oz.
Rosin	_	oz.
Rosin Oil	1	0Z.
Carnauba Wax	1	0Z.
No. 2		
Lactic Casein	10	oz.
Borax	2	oz.
	60	oz.
Pigment Sonn Solution	4	OZ.

0.5 oz.

0.8 oz.

remainder

Water remainder.

The soap solution can be sodium resinate or the potassium salt formed by boiling potassium carbonate (1 part) with carnauba wax (15 parts). The amonium sulphate is added after all the other ingredients are in solution. The pigment could be china clay or tale colored to shade with a brown lake. The composition given would require further adjustment with water to give the right consistency in the coating tank.

Numida Dyeing of Feathers
Dissolve gum arabic in cold water to
about the thickness of varnish.

Make up a solution containing:

Gum Arabic Water	1 glass
Cold Water	2 glasses
Glycerin	1 glass

Strain thoroughly to remove all particles of dirt, etc.

Take the dry feathers and work in this solution until thoroughly saturated, wring through the ordinary wash wringer, and squeeze out as much of the solution as possible, after which rub through the hands thoroughly for about 5 minutes in order to evenly distribute the remaining portion of the liquid that is in the feathers, after which string the feathers and beat them out on a wooden board for several minutes until the fine stems separate, after which hang up and dry overnight.

Feathers thus treated will retain this effect under all ordinary conditions.

Fabric Paint

I WOIT I WIND	
Basic Dye	2 lb.
Ethylene Glycol	60 lb.
Zine Chloride	6 lb.
Tunnic Acid	6 lb.
Glacial Acetic Acid	6 lb.
Gineral Acetic Acid	90 lb.
Tragacanth Solution (1%)	6 0 101

Synthetic Resin for Impregnating Textiles

British Patent 422,957

Polyvinyl Chloride (60- 65¢; Chlorine) Methylene Chloride Benzene Butyl Acetate	3	lb. lb. lb. lb.
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Weighting Cotton Yarn

Cotton yarn may be weighted to a considerable extent, when dyed with the direct colors, by adding magnesium sulphate (Epsom salt) to the dye bath, together with a small quantity of dextrin. Owing to danger of imperfections in the color, such as unevenness and cloudiness, it is perhaps better to use a separate bath after the dyeing for the purpose of weighting. This will be especially true if it is desired to weight to any considerable extent. The following process is a typical example of weighting cotton yarn which has been dyed with direct colors. For 100 lb. of cotton yarn use a bath containing about 160 gal, of water; add 100 lb. of magnesium sulphate, 15 lb. of dextrin, and 2 lb. of glycerol. Have the temperature of the bath at about 120° F. The cotton yarn is entered into this bath and turned for

20 minutes, or until the fiber is thoroughly saturated with the solution. It is then removed, hydro-extracted and dried. Such a treatment as this will give a weighting of about 10 to 12% to the cotton yarn. The bath is by no means exhausted, and may be freshened up by the addition of a small amount of magnesium sulphate and dextrin till it is brought back to the same hydrometer test as at first, and succeeding lots of cotton may be treated as above. The glycerol is added for the purpose of preventing the weighting material from giving the fiber a stiff handle.

Rayon Spinning Solution

To a solution of 25 parts acetonesoluble cellulose acetate and 75 parts of

95% acetone plus 5% water is added 2.5 parts of a mixture containing mineral oil (100 viscosity at 100° F. Saybolt) 85, saponifable oil (olive oil) 10, tetrahydronaphthalene 2.5 and soap (sodium oleate) 2.5%. The yarn spun from the solution is bright and fairly transparent and has superior knitting properties.

Wet Strength of Wet Fibers, as a Percentage of Their Dry Strength

Cotton	110-120%
Wool	80- 90%
Silk (True)	75- 85%
Acetate Silk	65- 70%
Cuprammonium Silk	50- 60%
Viscose Silk	45- 55%
Nitro Silk	30- 40%

MISCELLANEOUS

Boiler Compounds

Formula No. 1	
Sodium Alginate (Crude)	20 lb.
Extract, Quebracho	12 lb.
Soda Ash	10 lb.
Trisodium Phosphate	10 lb.
Caustic Soda	1 lb.
Water	300 lb.

Manipulation: Dissolve the salts in the water and add the alginate and quebracho extract at room temperature.

No 2 Anhudrana Disadium

Phosphate	47 lb.
Soda Ash	44 lb.
Corn Starch	9 lb.
T. 1 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1.2

It should be noted that this formula includes both inorganic and organic constituents. The starch is added to bring about a state of colloidal suspension of the insoluble matter precipitated in the boiler so that a sludge is formed in preference to a scale.

Another composition which deserves consideration is the U. S. Navy Standard Compound, which is:

No 3

	110. 0		
Anhydrous	Sodium Carbonate	76	lb.
Trisodium	Phosphate	10	lb.
Dextrin or			lb.
Cutch	sufficient to yield	2	lb.
	tannic acid		
Water	to make up to	100	lb.

Coal Dust Briquettes German Patent 616,376

Finely divided coal sludge brought to water content of 12 to 20% is mixed with 2 to 3% molasses and then compressed in molds and dried.

Fuel Briquettes for Motors

One hundred kilograms of sugar or molasses are mixed with 5 kg. of alum or a similar substance for inversion of the sugar and dissolved in 400 to 600 kg. of water, after which finely ground bitu-minous coal is added until a homogeneous mixture is obtained. The mixture is

poured over 50 to 100 kg. of a finely disintegrated mass of sugar beets. Thirty to 50 parts by weight of the mass thus obtained are mixed with 70 to 50 parts of finely ground charcoal, and the mixture is nney ground charcoal, and the mixture is pressed to briquettes under a pressure of 100 to 300 kg, per sq. cm. The briquettes are dried by heating in a separate drying chamber by means of combustion gases from a steam boiler furnace. The drying requires only about 15 to 30 minutes, during which the briquettes take on a coke-bkg anneagage. Onlying to the bright term. hke appearance. Owing to the high temperature in the drying chamber, about 350° to 500° C., and the high water content of the briquettes, steam is formed during the drying which seems to have a hardening effect upon the briquettes. Under this high drying temperature the sugar content of the briquettes is caramelized. A suitable composition of the dry mattereof the briquette mass is stated as 80 parts by weight of charcoal, 20 parts of bituminous coal, and 2 to 6 parts of sacchariferous binding substances.

Fuel Briquettes U. S. Patent 1,977,332

Slowly burning briquettes suitable for use in orchard heaters are formed by mixing chaicoal 50, sand 25 and a sugar-syrup binder about 25% so that all the particles of charcoal and sand are coated by the syrup, molding without applying pressure, evaporating moisture from the briquette in the mold and then heating to about 370° C. for about 2 hours to form an anhydrous porous mass, and cooling under air-tight conditions.

Briquettes

French Patent 766,979

Semicokes and fine coals are mixed with 6-12% of pitch, molded and heated to about 600° C. and then carbonized at 700-900° C.

Battery Paste

In the manufacture of lead-acid storage battery plates it frequently happens that the paste in the plates checks when dried. The addition of a small amount of silicate of soda to the paste will reduce this tendency. The amount should be not over 1 oz. of the strong solution of silicate of soda (water glass) to 100 lb. of the oxide. This should be dissolved in about 1 pt. of water and added to the oxide before the acid is added.

Low-Voltage Storage Battery Paste U. S. Patent 1,944,065

The paste for a lead accumulator contains (a) 0.9 to 1.5 weight per cent of nickel sulphate, or (b) 0.1 to 0.5 weight per cent of cobalt sulphate as active material.

Cold Storage Fluid

U. S. Patent 1,943,268

Fluids for cold storage comprise water (in each case) and butyl alcohol 10%, or ethyl ether of glycol acetate 20, or di-ethylene glycol butyl ether 5%.

Low Freezing Heat Transfer Medium U. S. Patent 1,972,847

A stable heat transfer medium comprises a mixture of 60 parts of diphenyl oxide, 12 parts of naphthalene, 28 diphenyl.

Antifreeze Composition

Formula No. 1

A mixture of 65% isopropyl and 35% methyl alcohol is recommended for addition to radiator water. It does not attack the metal parts and changes the boiling point of water only slightly.

No. 2

U. S. Patent 1,997,735

A cooling medium having a freezing point below -45° F. and a boiling point above 217° F. consists of a solution formed by adding 2 lb. of calcium chloride and 7 oz. of aluminum chloride to glycerin, 1 pt., and water as 1 gal.

Prevention of Ice Formation on Airplanes U. S. Patent 2.017.593

A mixture of liquids of different effects on rubber (such as pine oil 4, diethyl phthalate 4 and castor oil 1 part) is used in such relative proportions as not substantially to swell or otherwise deteriorate a rubber surface to which the composition is applied.

Anti-Knock Fuel

Formula No. 1

U. S. Patent 2.021.088

0.5 to 5% of ethylene diamine or 0.5 to 1% of a hydrate of the same is used with gasoline.

No. 2

U. S. Patent 1.973,320

A mixture for introduction into the cylinders of internal combustion engines to prevent knock or pinking and the deposition of carbon comprises 85 g. of uranium chloride and 15 g. of vanadium chloride dissolved in acetone.

U. S. Patent 1,980,097

Chloral hydrate in small quantities may be utilized to assist the solution of the metallic chlorides. For example, 1 to 10 mg. of platinum chloride may be dissolved in 1000 cc. of butyl oxalate and 250 to 2500 mg. of vanadium chloride in the same amount of butyl oxalate. The solutions are then combined and sufficient butyl phthalate is added until it consti-tutes about 25% of the mixture.

Stabilization of Anti-Knock Compounds British Patent 414,581

Decomposition of lead tetra-ethyl present in the fuels is prevented by the addition of a small amount, e.g., 0.01-0.05% of sodium fluoride, potassium fluoride or ammonium fluoride.

Detergent for Automobile Radiators Formula No. 1

U. S. Patent 1,967,393

A mixture is used comprising ammonium hydroxide or cyclohexanol 1, a sulphonic acid derivative of an alkylated aromatic hydrocarbon such as sodium 1-isoprioylnaphthalene-2-sulphonate about 0.4 and an alkali metal carbonate such as sodium carbonate about 4 parts.

No. 2

U. S. Patent 1,967,394

This relates to a detergent mixture comprising an organic solvent immiscible with water such as ammonium hydroxide or cyclohexanol about 1, a sulphonic acid derivative of an alkylated aromatic hydrocarbon about 0.4 and sodium phosphate about 4 parts.

Carbon Electrodes for Batteries British Patent 429,840

A mixture of finely-ground bone charcoal 34, wood charcoal 8, graphite 6, pinewood flour 8, ammonium sulphate 14, and sulphur 6 parts, with a binder made by stirring a mixture of wheat flour 6, sugar 18, water 7.5, and oil 15 parts at 80° C. for 15 minutes to burst the starch granules and dissolve the sugar, is extruded or pressed into the desired electrode shape, dried, and fired in cast iron boxes or in saggers packed in graphite or retort-gas carbon, the temperature being raised slowly to 1000° in 16 hours and maintained there for 4 hours. After cooling, 1/4 of the block is immersed in a 2-5% solution of paraffin wax in petrol and the other % is then immersed for 3-4 minutes in 10% aqueous ammonium chloride. The waxed top is then drilled, a copper terminal screwed in, and the joint again waxed. Finally the whole electrode is impregnated with a 10-12.5% solution of silicic acid in trichloroethylene, carbon tetrachloride or other volatile solvent and dried.

Brake Fluid Composition U. S. Patent 1,928,956

A hydraulic fluid comprises, in solution, glycol acetate, e.g., 50% by volume, with smaller proportions of water 37-45 and sulphonated castor or linseed oil soap, 5-13.

Moisture-Resistant Bristles U. S. Patent 1,953,980

The bristles are first impregnated with an aqueous heavy-metal sult (e.g., 1-3% aqueous aluminum acetate) and then with a water soluble soap of a fatty acid (e.g., 4% aqueous castile soap). They may also be dipped into a solution of a wax in xylene.

Catalyst Canadian Patent 350,894

To a dry mixture of kieselguhr 150, gum tragacanth 10 and potassium sulphate 20 lb. is added with agitation a sodium vanadate solution prepared by treating 16 lb. of vanadium pentoxide with 10 gal. of water containing 11.3 lb. of sodium hydroxide. The mixture is diluted with 20 gal. of water and after thorough mixing sulphuric acid is added to neutralize or nearly neutralize the mixture. The mixture is evaporated to a consistency suitable to permit granula-

tion or pelleting and the granules or pellets are heated for 1 hour at 600° C. The product is a catalyst for the oxidation of sulphur dioxide.

Catalyst for Ammonia Oxidation U. S. Patent 2.017.683

Metallic colult, containing impurities 70, is heated to effect fusion with calcum carbonate 3.5-5 and calcium fluoride 1.7-3.5 parts, the slag formed is separated from the metal and the latter is converted into colult oxide.

Activation of Kaolin for Catalytic Purposes

Kaolin is ignited at 750-800° C. for 2 to 3 hours and treated in the cold with 33% intric acid for 24 hours and the solution is then heated at 60-80° C. for 3 to 4 hours. Aluminum hydroxide is then precipitated and allowed to stand for 1 day at room temperature before filtration. It is dried at 100 to 120° C. and activated at 360-385° C. The actulyst is suitable for the dehydration of alcohol.

Regeneration of Spent Nickel Catalysts

The method consists essentially in treating the spent catalyst successively with a small quantity of 20° Bé. sodium hydroxide, sulphuric acid and water. Before suponifying the spent catalytic mass, it is heated with indirect steam with vigorous stirring till a homogeneous mass is obtained. The sodium hydroxide solution (60-80 l. for 500 kg. of catalyst) is then added, followed by sufficient water to make the mass fluid; saponification is effected by heating with stirring for 11/2 to 2 hours. After transferring the soap to a lead lined tank, it is decomposed with concentrated sulphuric acid, diluted with water and allowed to stand, and the supernatant fat is removed. The nickel is then boiled with sulphuric acid as usual. The recovery of nickel is 92-94%, as compared with 64-70% by the ordinary method.

Fuel Catalyst * French Patent 765,824

A mixture used for activating the combustion of solid fuels contains, e.g., manganese dioxide 32.1, organic material (wood charcoal) 2.5, sodium chloride 27.7 and sodium chlorate 37.7%.

Cable Insulation U. S. Patent 1.946.322

The mixture comprises a hydrocarbon oil (e.g., cylinder oil) 95-50, and rosin free from oxidized components, especially abietic acid. 5-50%.

U. S. Mint Test Solutions for Counterfeit Coins

Concentrated Nitric

Acid 61/2 drachms Hydrochloric Acid 15 drops Distilled Water 5 drachms

Silver

Silver Nitrate 24 gr. 30 drops Nitrie Acid Distilled Water

A drop of the above solutions will have no effect on genuine coins; but will stain others, i.e., spot them.

1 07.

Capsules British Patent 412,975

Capsules or coverings, for bottles, jars, metal tubes and rods, of the kind made from a composition containing cellulose ester and a substance which may be removed by a suitable solvent after formation of the capsule, etc., to cause the capsule, etc., to shrink on drying and fit tightly onto the article to which it is applied, are formed by compression, extrusion or injection from a composition produced by working or mixing together the cellulose ester, a water soluble softener and optionally, a phasticizer to solid but plastic composition. Small amounts of a volatile solvent may be amounts of a volatile solvent may be and optionally, a plasticizer to produce a example, a mixture mixing, in an example, a mixture containing cellulose acctate 3, monochlorohydrin 2, monoglycrol benzoate 1 and water 2 parts is mixed at 80-100° C, until completely gelatinized and most of the water has evapo rated. The material is then formed to the desired shape and rendered contractile by soaking in water to dissolve out the monochlorohydrin. The contractile capsule is then applied to the article on which it is to be used and, as the water dries out, the capsule shrinks into posi-tion. Filling materials, dyes or pigments may be added.

Motor Carbon Remover U. S. Patent 2,004,628

A carbon removing composition is composed of kerosene, creosote, castor oil and

amyl acetate, combined in substantially the following proportions: kerosene, 491/2%; creosote, 25%; castor oil, 25% and amyl acetate, 1/2%.

Activating Adsorbent Clay U. S. Patent 1,976,127

The method of activating adsorbent earths comprises mixing an earth with concentrated sulphuric acid in an amount equal to from 5% to 35% of the weight of the earth, heating the mixture to a temperature of 150 and 300° C. to obtain reaction of sulphuric acid with constituents of the earth and to also partially dry the earth and the products of such reaction by the combined effect of heating and the dehydrating action of the sulphuric acid, then bringing the resultant mixture into contact with water to dissolve soluble salts therefrom, separating the solution from the undissolved earth, and then drying the earth.

Processing Coal Canadian Patent 324,976

Coal containing iron sulphide is thorroughly washed to remove dust and impurities and while wet is sprayed with a compound containing calcium chloride 92, potassium dichromate 3, manganese dioxide 3 and tannic acid 2 parts by weight. The burning properties and ash characteristics are improved and the deleterious effect of flue gases and tube-slagging is minimized.

Oil Treatment of Coal U. S. Patent 2,005,512

The process of treating solid lump fucl to render the same dustless, consists of heating oil having a gravity of 10° to 30° Bé. at 60° F. and a Saybolt viscosity of 100 to 1200 at 100° F. to a spraying temperature of 100° to 250° F., and spraying the heated oil in finely atomized state on the fuel in quantities sufficient to deposit on the fuel a thin enveloping film of oil.

Fuel Oil Activator Japanese Patent 101.701

Naphthalene Anthracene Phenanthrene

100 oz. 5- 10 oz. 1- 3 oz.

Thirty grams of the above is added to 5 gal. fuel oil to increase heating efficiency.

Dustproofing of Coke

A 1 to 1 emulsion of a thick petroleum oil and water is made at 94° C., and then diluted with 7 parts of water at 38° C. Two gallons are sprayed per ton of coke on the loading chutes.

Decolorizing Charcoal from Corncobs Soak corncobs in 3% zinc chloride and 7% sulphuric acid for 24 hours. Distill destructively at 600° C. for 50 minutes and treat with superheated steam at 400° C.

Deodorizing Petroleum

Petroleum products may be conveniently deodorized by agitating thoroughly with quicklime, 3 oz. to the gal. and filtering.

Gasoline Gum Inhibitor U. S. Patent 1,970,339

Nicotine pyrogallate or amylgallate is added in proportion of about 1/100%.

Coloring Leaded Gasoline Canadian Patent 352,875

a(2 Methoxyphenylazo)-2-naphthol is used at rate of 2 to 12 oz. per 10,000 gal.

Liquid Dielectric Composition

U. S. Patent 1,999,004

Chlorinated hiphenyl having a chlorine content of 60% is used in a proportion of 45% together with trichlorobenzene 25 and tetrachlornaphthalene about 30%.

Condenser Dielectric

A 50% solution of Bakelite in castor oil has a high dielectric constant, 5.6, as compared with 2.3 for transformer oil. A condenser having paper impregnated with the Bakelite mixture has a power factor of 1% against 0.5% with transformer oil, this being the only disadvantage.

Dielectric Materials

French Patent 765,876
Dispersions of metal soaps in insulating oils are used, e.g., 6-10 g. of aluminum stearate in 94-90% of oil.

"Coreth" Type Artificial	Diesel Fuel
Alcohol	36 kg.
Coal-Tar Oil	28 kg.
Gas Oil	20 kg.
Wood Oil	10 kg.
Water	4 kg.
Degras, Saponified	2 kg.

Liquid Electric Insulation British Patent 413.596

A mixture comprising mineral hydrocarbon oi (50-70 parts) and halogenated diphenyl, e.g., the polychlorinated derivative (50-30 parts).

Electrical Insulator British Patent 429,730

Rutile	32	lh,
Talc	58	lb,
Blue Clay	6.5	lb,
Calcium Carbonate	3.5	lb.
Mix thoroughly and mold.		

Electric Insulating Compositions German Patent 616,056

A binder for use in making insulating compounds or maternals comprises a resin, a vegetable drying oil, shelline and (as a flux) an aromatic compound boiling above 200° C. A specified binder comprises copal 12.5, wood oil 1.5, α-nitronaphthalene 1 and shelline 10 parts. Mixtures of the binder with subdivided mica of like material may be molded under heat and pressure, or a solution of the binder in an organic solvent may be applied to mica sheets and the latter then united by heat and pressure.

Electrical Insulating Fused Magnesia British Patent 413,905

The electrical resistivity of fused magnesium oxide is permanently increased by heating slowly to 1149° F. maintaining it at this temperature for about 6 hours and finally cooling to room temperature in about 30-40 hours

Vitreous (Electrical Insulating) Material

U. S. Patent 1,984,178

Silicon dioxide is fused with beryllium oxide 0.14-1.5 and aluminum oxide 0.2-2.0%.

Waterproofing Electrical Wires

Formula No. 1	4
Crepe Rubber	\$ 0 lb.
Mineral Spirits	30 lb.
Mill together until unifor while mixing	m, then add
Glue Solution (20%) followed by	25 lb.

20 lb.

p					
	No. 2				
U. S. Pa	tent	1,963	,895		
Mineral Oil					cc.
Neat's Foot Oil Ethyl Acetate					cc.
Emyr Acetate				•	
Electrical 1	[nsula	ting	Taj	96	
Form	ula N	lo. 1			
Make up caoutel			ions		
a. Caoutchouc, C Smoked Pres	Crude	, in		20	l
b. Benzene or I	Benzo	line	•	80	
	Vo. 2				•
Resin C	ompo	sitio	ns		
Formul		b	0	ď	
Rosin Rosin Oil	40 36	30 30	20 28	20 30	
Rosin Tar		10			g. g.
Petroleum Tar		_	10	_	g.
Stearin Tar Coal Tar	_	_	10	20	g. g.
Wood Tar,					٥.
Anhydrous Mineral Oil	_	10	-8	10 10	g.
Linseed Oil	24	20	24	10	g. g.
The formulae a	and	b ar	e su	peri	or to
the two others. F pale resins, as a, a	or w	liite ssibl	ribb o	ons,	only
	Vo. 3	00101	·.		
Fillers		Pigm	ents		
For White Ribbon			a	b	
Lithopone Zinc White			80 20	60 20	g.
Barium Sulphate	1			20	g. g.
For Black Ribbons	3		c	d	Ü
Barium Sulphate Vegetable Black	1		40 45	20	g.
Lamp Black			15	15	g. g.
Frankfort Black			-	45	g.
Chalk Powder	To. 4			20	g.
Definitive Mixture					
White Cover Ribbo	n:				
Rubber Solution Resin Composition	(No. n (N	1) [0. %	7)	38 22	g.
Fillers (No. 3a,	36)		•,	40	g. g.
Black Ribbon:	/3T	• • •	a	b	
Rubber Solution Resins (No. 2b.	(2c)	1)	44 26	42	g. g.
Regine#(No. 9d)	-			30	g.
Fillers (No. 3c,	3d)		30	28	g.
W .	o. 5 Be A	nnlie	d b	.	
Coating to Be Applied by Hot Impregnation					
White Coating					
a. Linseed Oil (Crude Rubber	60°C	'.) Smal	,	57	g.
Piecer	,	omqı	•	6	g.
					-

b. Resin	9 g.
c. Fillers (No. 3a, 3b)	28 g.
Prepare a in a kneading	machine at
50° C., then heat up to 180	
clear solution is formed, add	d melted b.
Cool to 100° C., and add o, st	ir, and dis-
harma abona 700 C	,

Black Coating		
Linseed Oil	60	g.
Crude Rubber	6	g.
Rosin Tar	12	g.
Fillers (No. 3c, 3d)	22	g.
or		
Mineral Oil	52	g.
Crude Rubber		g.
Petroleum Tar	12	g.
Wood Tar, Anhydrous	10	g.
Fillers (No. 3c, 3d)	20	
These masses should be kept	at	60-

80° C. in the impregnation vat.

Fusible Cut-Outs British Patent 423,076

A fuse wire incorporated in a current consuming device, e.g., an incandescent or arc-discharge lamp, rectifier or valve, is composed of a brass containing 0.25 to 8% aluminum, e.g., copper 67, zinc 32, and aluminum 1%. This is non-oxidizable, has a higher resistance and melts rapidly.

Electrolytic Condensers British Patent 421,628

An electrolytic condenser is wound in annular form to permit free circulation of air around it. Aluminum electrodo strips are separated by strips of cloth impregnated with an electrolyte, of the composition glycol 400 cc., borax 25.6 oz., boric acid 17.0 oz. and water 25.6 oz.

Electrolytic Condenser Medium U. S. Patent 1,973,554

Monoethanolamine	1 lb.
Ethylene Glycol	5 lb.
Boric Acid	5 lb.
Heat together until dissolu	hha haa ha

until dissolved and add bentonite or starch to consistency desired.

Fingerprint "Raising" from Cloth

Dip in, or paint with a 10% solution of silver nitrate to which has been added a little acetic acid. Dry in dark room, then expose to ultra-violet light until of maximum. mum intensity, and photograph.

Latent Fingerprinting

A piece of paper or other material on which one is searching for fingerprints is saturated in a sensitizing solution prepared by dissolving 2 g. of silver nitrate in 1 l. of distilled water. This is stored in a dark place. After having soaked for 2 hours in the silver nitrate bath, the paper is thoroughly washed in distilled water, first by soaking for 30 minutes and then two rinsings. There is left in the paper only the silver chloride which has been formed from the chlorides left by the perspiration and the silver nitrate. The paper is hung up and allowed to dry thoroughly. It is then developed, either with a developer of the M. G. type or with others, such as formaldehyde and sodium carbonate. Following the development the paper is again washed in water, then in a bath of hypo, washed, and dried, and is ready for observation.

If kept in a humid atmosphere the migration of the chlorides may be so intensified that in time a gray cloud is formed where the print was originally. In some cases the print goes through the paper. Prints made from the skin of a corpse are very poor and diffuse, although chloride is deposited.

Fire Extinguisher U. S. Patent 2,010,729

A fire extinguishing composition comprises 48 parts by weight of sodium bicarbonate, 12 parts by weight of horic acid, 41/2 parts by weight of potassium bitartrate, and about 11/2 parts by weight of borax.

Fireproof Film Containers British Patent 419,249

The walls are made of a mixture of sawdust 25, calcined magnesite 25, magnesium chloride (as a 25% aqueous solution) 30, potassium alum 10 and a mixture consisting of asbestos flour 4, asbestos fiber 3 and acetic acid 3, 10 parts.

Fluorescent Screens French Patent 770,728

The screen contains zinc or cadmium borate, e.g., zinc silicate 10-12, calcium 45-50 and zinc or cadmium tungstate borate 40-45%.

Electrotyping Matrix British Patent 430,660

A sheet of aluminum 0.007 in. thick, is cleaned with etching fluid or caustic soda and then coated with molten beeswax, preferably a mixture of gnm damar 2 and beeswax 16 parts, heated to 160° F. The wax face is then coated with graphite to render the surface conductive,

Masking Taste of Chlorinated Water Add 2 or 3 tablespoonfuls of wine to each liter of water.

Fish Baits

The common "baits" comprise two general cutegories: (1) artificial baits, and (2) natural baits.

Artificial buts muy be classified as flies, spoons, sponners, phantoms, and a multitude of other contrivances, some of which may be used alone, and some in combination with natural barts. Flies are largely unde of feathers, worsted, silk, tinsel, etc., and are fashioned to suggest an insect. Most flies, however, resemble only remotely any known insect. Other barts may be made of metal, wood, rubber, etc. The list is too extensive to enumerate here, and more may be learned from a reliable fishing tackle dealer than from reading pages of descriptions. With reference to natural buits, with which the following lists are concerned, a local nugler can usually impart to the novice more practical knowledge in a short time than could be learned from a whole volume of discussion and descriptive matter.

Judging by the stomach contents of fishes, there are but few groups of animals, from werms to mammuls, that do not afford food for one or another game fish. That some of these are occasionally swallowed by a fish, however, does not necessarily signify that they would make good bait. Furthermore, some of the best baits can never be the natural food of the fish. The groups of animals which comprise forms most commonly employed as bait, from the lowest form up, are: worms, mollusks, insects, crustaceans, fishes, birds, and mammals.

It must be borne in mind that baits used in one part of the country may be of little or no avail in another part, even for the same species of sish; and that in the same locality the preper baits often vary with the time of mean. Fur-thermore, a killing bait of one my may prove ineffective on the next. Success in fishing, therefore, depends largely upon the experience, judgment, skill, and patience of the fisherman.

Vernacular names of the various animals used for bait differ greatly in different parts of the country; for instance, the stone fly of one section is the mill fly of another, and the hellgramite of one locality is the dobson of another, and so on. Therefore, any list of baits can be of only partial assistance. The following lists aim to give the most common baits under the names by which they are most widely known.

Natural Baits are used in several different ways, such as in still fishing, bait casting, skittering (modified form of casting), or trolling.

Live Bait .- It has always proved practically impossible to keep a large amount of live bait in restricted limits; furthermore, no flah will live indefinitely without food. The kind of food necessarily depends upon the kind of fish, but most shiners and other minnows are more or less carnivorous and finely ground ment of some kind would probably answer for this class. The most appropriate food, however, would be small crustaceans and aquatic insects such as are usually present in sluggish streams and small ponds. These may be collected by means of a gauze dip net. It is possible to stock a small pond or pool, or even a rain barrel, with small crustaceans and maintain a supply of that kind of food. Some species of bait minnows are much hardier than others, but in all cases, when kept in confinement much depends upon the maintenance of cleanliness and a sufficient supply of oxygen. The needed oxygen is best supplied by a continuous flow of well acrated water, but where this is impossible it may be furnished in a fine spray of compressed air introduced near the bottom of the tank. Cold water will dissolve more oxygen than warm water, therefore, the temperature should be kept low if possible. Overcrowding should be carefully avoided and all injured or sick fish should be removed as soon as detected. If feeding should be attempted great care should be taken to remove all food uneaten, as otherwise it will decay and pollute the water.

Conditions will vary according to the species of ninnow, the size and character of the tanks or pools, the temperature of the water, and the number of fish per unit of space, and it is difficult, therefore, to furnish specific information without a knowledge of these factors.

Keeping and Rearing Earthworms for Bait.—Earthworms multiply by producing eggs which are laid in capsules in the ground. The young become fully grown in four or five months. One method of culture is to sink into the soil

in some shady spot a box of suitable size, usually not less than 18 inches deep and of any desirable width. The top of the box should be made hinged, or removable, and placed from two to three inches below the surface of the surrounding soil. This box should be nearly filled with rich, dark loam that should be kept quite moist but not wet, for too much water will kill the earthworms quickly. The worms may then be collected and placed in this box, and may or may not be covered with a layer of green sod.

By far the easiest and most convenient way to collect earthworms is by the use of a flashlight or lantern at night. They may be found on the surface of ground which has been devoted for some years to lawn or sod purposes. The worms are usually much more numerous during the months of April, May, and June than at any other time, although they may be easily brought to the surface at any season of the year, except winter, by thoroughly sprinkling the soil in the early evening. If food is provided for the worms in the box, they may be kept almost indefinitely in such container without changing the soil. They have been raised successfully by feeding ordinary molasses spread on one side of a gunny sack, which is then laid on the surface of the ground with the sticky side downward, and the back of the bag then sprinkled with water. Powdered bread crumbs and crumbled hard boiled eggs have also been used as food.

Fresh Water Crawfish and Shrimp, Keeping Them Alive for Bait.—These crustaceans can be kept alive in tanks, simil pools, or wooden boxes which are well supplied with running water. The best food for them is fresh meat fed in small pieces, but great care should be taken not to leave old and spoiled meat in the water for any length of time, as this will soon prove fatal. The boxes or other containers should not be over-crowded and should be cleaned often and the dead crawfish or shrimp thrown out, as they decay rapidly and will soon cause the death of the healthy ones. The same general treatment is used if the crustaceans are to be kept in closed tanks or aquaris.

Hellgramites.—These are the larval form of the dobson fly. They are found under stones in swift streams and are an excellent bait for bass. Hellgramites can be kept alive for a considerable time in floating bait boxes or in wet grass.

Glow Worms .- The term glow worm is

applied to the wingless female beetles of the family Lampyridac. They are They are nocturnal in habit and feed upon smaller insects and worms. They can be kept alive in loose, damp earth, covered with moist grass and kept in a cool place.

Preserving Minnows for Bait .- Take 1 part of formalin to 29 parts of water, place the minnows in this solution in a tightly closed jar or bottle and keep in the dark until they are to be used. In this way they will retain their colors and silvery hues better than if in the light.

When about to use the bait, soak it in fresh water to remove the formalin. A few drops of oil of rhodium may then be placed on the minnow to disguise the pungent odor of formalin that may remain in the fish after soaking. The oil of rhodium is said to be attractive to fish but be that as it may it does not repel them as the formalin is likely to do.

Dough Balls .- A tough paste may be made of moistened bean, wheat, or other flour, thoroughly mixed with a little sugar, or preferably honey. To give the paste a greater tenacity, cotton batting or wool should be stirred in. Ground or mashed white meat, such as veal or pork, or any bleached meat may be added, but this bait must be fresh and kept untainted. Dough balls may be made also by boiling rye flour to a consistency of paste, then sprinkling with corn flour and rolling into a "ball."

List of Common Fresh Water Game Fishes with General Mention of Some Baits Used in Their Capture

Bowfin, Dogfish, Grindle

Frogs, minnows, pieces of fish, etc. Blue Cat, Chuckle-Headed Cat, Fulton Cat

Minnows, shiners, worms, crawfish, pieces of fish, meat, liver.

Spotted Catfish, Channel Cat, Fiddler Shiners, worms, meat, liver, dough

balls.

Common Bullhead, Brown Bullhead, Speckled Bullhead

Minnows, worms, frogs, grasshop-pers, pieces of fish (chub, perch, sun-fish), salt, mackerel, salt pork, meat, liver.

Mud Cat, Yellow Cat, Goujon, Morgan

Crawfishes, fresh hickory shad, other fish baits. Buffalo Fish

Worms, insects.

Carp Sucker Worms, insects.

Sucker

Earthworms, bits of crawfish.

Redhorse

Worms, insects. Chub

Pieces of fish, insects, grasshoppers, Worms.

Squawfish

Worms, minnows, shiners.

German Carp

For angling, various baits have been recommended. Worms, grubs, grass-hoppers, and pieces of fresh meat have been used successfully, but the most highly recommended baits are composite pastes. Pellets of partly boiled potatoes are said to be good bait, as well as dough balls or corn kernels wrapped in mosquito bar.

American Ecl, Fresh Water Ecl Earthworms, shiners, grasshoppers, etc.

Mooneye

Minnows, worms, msects.

Common Whitefish

Worms, insect larvae, may flies, shrimp, pieces of fish, minnows.

Rocky Mountain Whitefish

Worms, insects, fresh ment.

Salmon, Sca Salmon, Eastern Salmon Worms, smelt, shiners, pork rind.

Landlocked Salmon Smelts, shiners, worms.

Black Spotted Trout, Cut Throat Trou Worms, grasshoppers, insects, minnows, pieces of meat.

Steel Hend Trout

insects, grass-Shiners, worms. hoppers.

Rainbow Trout

Worms, grasshoppers, insects, shiners. Brown Trout

Worms, various insects, grasshoppers, crickets, shiners, minnows, pieces of fish, horse meat.

Lock Leven Trout

Worms, various insects.

Chinook Salmon

Smelts, shiners. Brook Trout

Earthworms or "barnyard hackle," grasshoppers, grubs, crickets, beetles, bumblebees, caterpillars, mill fly, caddis fly larvae, may fly, newts, mice, or bits of animal flesh. A capital bait is the beautifully tinted anal fin of a trout, which in water with some current waves wabbles and flutters in a most seductive manner on the hook.

White Trout, Golden Trout Worms, pieces of fish, smelts, minnows, shiners.

Dolly Varden Trout

Worms, minnows, shiners, insects.

Lake Trout Minnows, shiners, pieces of fish (Whitefish), ciscoes.

Grayling

Gaddis fly larvae, "rock worm," earthworms, grubs, crickets, grass-hoppers, natural flies, or small bits of fat meat.

Smelt

Pieces of smelt, shiners, minnows, worms, shrimp. Common Pike, Pickerel

Frogs, shiners, minnows, white chub, pork rind, fish belly, 3-4 in. piece pickerel stomach, perch belly.

Muskellunge Small fishes, suckers, shiners, ciscoes, grasshoppers, frogs. White Crappie

Worms, minnows, shiners.

Black Crappie, Calico Bass Minnows, worms, small shiners.

Rock Bass, Redeye, Goggle-Eye Small minnows, white grubs, earth-worms, grasshoppers, crickets, small crawfish, yellow perch, fresh water mussel, frogs.

Warmouth Bass

(See Rock Bass.)

Red Robin, Long Eared Sunfish Worms, insects, minnows. Bluegill, Blue Sunfish

Worms, insects, insect larvae. shrimps, small crawfish, pieces of fresh water mussel.

Green Sunfish, Blue Spotted Sunfish Worms, insects, insect larvae. Pumpkinseed

Worms, insects, pieces of crawfish, pieces of meat.

Shell Cracker

Worms, insects, small crawfish, pieces of fish.

Black Bass

The best natural bait is the minnow, a shiner, chub, or the young of almost any fish, which is well adapted for either casting, trolling, or still fishing. In waters where it abounds, the crawfish is a good bait, especially the shedders or soft craws, to be used only for still fishing. The hellgramite, the larva of the corydalis fly, in its native waters, is also successful for still fishing. A small frog is capital bait a weedy waters, where it is usually cast overhead with a very short and stiff rod. Grasshoppers and crickets are sometimes employed with a fly-rod, in lieu of artificial flies, with good results. The salt water shrimp, where it is available, near the coasts, is also a good bait for still fishing. Cut bait

is also sometimes useful. It should be remembered that all baits of whatever kind, should be kept in motion. A dead minnow answers as well as a live one for casting or trolling, but should be alive for still With crawfish, worms, or hellgramites, a float fishing. shrimps or hellgramites, should be employed to keep them from touching the bottom. In casting the minnow it should be hooked through the lips, and reeled in slowly after each cast to imitate the motions of a live one as much as possible.

Large Mouth Black Bass, Oswego Bass Live minnows and other live baits, such as grasshoppers, frogs, hellgramites, efts, worms.

Small Mouth Black Bass Shiners, chub, small yellow perch with dorsal fin cut off, mad-tom, stone catfish, floor of mouth of pickeral cut like a fish, belly of bowfin, crawfish, hellgramites, crickets, efts, newts, small frogs, worms. Wall Eyed Pike, Pike Perch, Jack

Salmon Live minnows, as fallfish or dace,

corporal, roach, redfin, gudgeon, brook chub, piece of fish, worms. Yellow Perch, Ringed Perch, American Perch

Worms, minnows, crickets, grasshoppers and other insects, small fishes, small frogs, crawfish, pieces of fish. Striped Bass, Rockfish

Shiners, minnows, pieces of fish.

White Bass

Live minnows, grubs, earthworms. Yellow Bass

Minnows (live bait), worms.

White Perch Worms, grasshoppers, insects, min-

Fresh Water Drum, Croaker, Sheepshead, White Perch

Crawfish, pieces of fish, mollusks. Burbot, Ling, Eel Pout, Cusk

Yellow perch, sunfish, lamprey, crawfishes, pieces of fish, smelts.

Cut Flower Vitalizer U. S. Patent 1.978,201

Eight ounces of sugar or saccharin, 2 oz. of kaolin, 1 oz. of yeast, ½ oz. of charcoal, 1 cc. of oil of pine and ½ oz. of lime. The foregoing makes up a composition weighing about 12 oz. and this may be dissolved in a suitable amount of water. It has been found in practice that this diluted solution shows benefit to all cut flowers.

The benefit is so decisive that increased turgidity and intensified color in the tissue of leaf and petal are visible to the eve usually within 30 minutes after the flower stem is immersed in the diluted solution. This increased turgidity and intensified color is retained by the flower whether under average room temperature of 70° F. or in refrigerated temperatures of 40-50° F., although a cooler temperature, as when untreated, prolongs the life of the cut flower.

The treated cut flower under observation slowly continues its development, retaining a healthy and nourished appearance, to eventually produce seed as large and apparently vital as it would upon the parent plant which had been un-

usually well cared for.

Furthermore, a flower cut in the bud develops normally when treated in this solution; for instance the chrysanthemum cut when the bud first shows color will develop into a flower equal in every respect to its companions left uncut on the greenhouse bench.

Again the treated flower lasts much longer after being removed from water, in treatment such as florists must subject flowers to in funeral pieces.

Preserving Foliage

A method of preserving foliage consists in placing the leaves in a solution of glycerin 1 part, water 9 parts. The leaves are then removed from this solution, dried between blotting paper and pressed.

Gas Mask for Sulphur Dioxide

Fixtnel nose-bag masks, 7 in. by 8 in. and held over the face by rubber bands, are used as a protection against sulphur dioxide gas. Masks are soaked in the following solution:

Distillad Water 1000 cc.

Distilled Water 1000 cc.

Glycerin 250 cc.

Soda Ash 200 g.

Masks are worn while wet with the

Gas-Producing Material for Inflating Hollow Rubber Articles British Patent 416,591

solution.

A mixture of sodium nitrite 56.5, ammonium chloride 43.5, and ammonium carbonate 10 parts is inserted into hollow rubber articles prior to vulcanization; on heating carbon dioxide, ammonia and nitrogen are evolved which ex-

pand the article up to the mold during vulcanization.

Manufacture of Luminescent Materials British Patent 414,905

A 2 to 1 mixture of zinc oxide (or magnesium oxide) and germanium dioxide is moistened with dilute aqueous manganese chloride and sintered at 1000° F, to produce zinc (or magnesium) germanate, which fluoresces bright greenishyellow (or orange scarlet) under excitation with cathode rays.

Match Striking Surfaces British Patent 411,688

An ignition surface, suitable for self-lighting eignrettes, etc., comprises a mixture of amorphous phosphorus and a collulose derivative binder of the character of cellulose acetate. A mixture of 4 g, of amorphous phosphorus in 25 cc. of a 5% acetone solution of cellulose acetate is spread as a film on a suitable surface. Other solvents, e.g., ethyl acetate, may be used.

Microscope Slide Cleaner

Xvlol	1	fl.	OZ,
n Butyl Alcohol	1	Ħ.	UZ.
Alcohol, Anhydrous	2	A.	OZ.
Water	1	fl.	OZ.

Sterile Modelling Clay U. S. Patent 1,979,016

Seventy grains of chlorthymol for every 100 lb. of manufactured modeling clay are sufficient to render the same sterile and to preserve its hygienic condition for long periods of time. The finished product may be packed in airtight containers for shipment and storage to prevent possible oxidation of its ingredients.

Preserving Fluid for Museum Specimens

12-25	oz.
10	0 z.
0.1	07.
to make 100	oz.
	10 0.1

Removing Formaldehyde Odor from Museum Specimens

Wash with water and submerge for 1/2 hour in:

Urea				5	OZ.	
Ammonium	Phos	phate		1	0Z.	
Water				94	0Z.	
If the spe	cimen	is to	be	replac	ed	ip

formaldehyde it should be washed free of urea.

Colored Neon Lights U. S. Patent 1,951,006

A mixture of approximately 10% of argon with 90% of neon emits a lavender colored light. The proportions of the gases may vary widely, the colors and shades changing with the different compositions. It is preferable to employ from 5 to 25% of argon, the balance be-ing principally neon. The addition of carbon dioxide to the mixture of neon and argon, for example, results in a white or substantially colorless light. Therefore, introduce a substance such as calcium or magnesium carbonate, which is capable of releasing carbon dioxide to the tube containing rare gases such as neon and argon. When the tube is energized, carbon dioxide is released, and produces the white or substantially colorless light until the modifying agent is exhausted. Such tubes have been operated for more than 700 hours without change of the light emitted.

In introducing the modifying agent to the tube, several methods may be employed: The agent may be supported inside the electrode; it may be attached to the electrode; it may be coated on the wall of the tube or electrode chamber; or it may be simply deposited in the electrode chamber or in the path of the discharge through the tube.

Other modifying agents may be used, for example, a suitable hydride such as magnesium hydride can be used to maintain a trace of hydrogen in the tube in admixture with the gases therein to effect a desired change in the color of the light emitted when the tube is energized.

Electrode, Neon U. S. Patent 1,926,336

The electrode comprises a compressed cylinder of an intimate mixture of tantalum carbide (88%) and cesium chloride, rubidium chloride and lithium chloride (12%).

Oxalie Acid from Corncobs

Corncobs 100 lb. Nitric Acid (95%) 3 lb. Heat until dissolution is complete; cool

and add: Nitric Acid (50-55%) 3 lb.

0.1 lb. Vandium Pentoxide Allow to stand for 2 or 3 days; filter

and evaporate the filtrate to obtain crude oxalic acid which may be purified by recrystallization.

Radiator Corrosion Inhibitor U. S. Patent, 1,992,689

For preventing corrosion in motor radiators containing alcohol and water the following formula is used:

a. Triethanolamine a. Triethanolamine Phos-	0.33	0Z,
l phate	1.50	oz.
b. Triethanolamine	0.75	oz,
Lard Oil	0.75	oz.

Mix ingredients of b and stir into a. The above is used per 100 parts of al-

Scale Preventing Mixture

Formula No. 1

French Patent 776,235

A mixture of formic acid 100 and digallic acid 6 parts is used.

No. 2 French Patent 776,234

A mixture of digallic acid 100, and trisodium phosphate 60 parts, is used to prevent scale in motor car radiators.

Non-Corrosive Chlorinated Solvents

U. S. Patent 1,966,881

Five-tenths to 2% of pinene is added to prevent corrosion.

Tellurium Alloy Rectifier U. S. Patent 1,961,825

The rectifier consists of plates of magnesium and an alloy of:

Tellurium	97.5	oz.
Copper	2	oz.
Silver	2.5	oz.
Sodium	0.5	oz.
Adala and mall to the state of		

which are welded together by passing a current from one to the other with a film of water between them.

Aluminum Reflector Etching U. S. Patent 1,999,042

Using hydrofluoric acid and nitric acid the aluminum is first dipped into a solution of 1 part concentrated hydrofluoric acid in 19 parts of water at a tempera-ture of 50 to 60° C., until an etch of the desired depth is obtained. The surface is washed and the article is im-

mersed for several seconds in a solution of nitric acid containing 1 volume of acid to 1 volume of water and held at room temperature. The aluminum is washed and dried and a clean, bright and uniformly etched surface is obtained. In the sodium hydroxide-sodium fluoride etching procedure a 5% sodium hydroxide solution in water containing about 4% sodium fluoride is used. The aluminum is immersed in this solution at a temperature of about 90° C. until the desired etch is obtained. It is then removed, washed, and treated with a 1:1 nitric acid solution, washed and dried as before. Again a very satisfactory clean, bright, and uniformly etched surface is obtained. It should be noted that the presence of copper in the aluminum causes the metal to turn gray to black on immersion in the hydrofluoric acid or the sodium hydroxide solutions. This black coloration, due to copper, is removed by immersion in the nitric acid. However, the nitric acid does not remove the gray film due to graphitic silicon, if it is present, and this must either be removed by rubbing or prevented from

The effect of the presence of a sufficient amount of copper in aluminum on its etching properties is pointed out specifically by the following examples: A sample of a commercial grade of aluminum containing about 1% of impurties, including 0.6% iron, 0.3% silicon, and 0.01% copper, when etched with hydrofluorie and nitric acids as above described, produces a surface which is irregularly etched, having a streaked appearance. On the other hand, a sample of aluminum containing 0.6% iron, 0.08% copper, and 0.18% silicon as impurities, when etched in a similar manner, produces a very satisfactory uniform reflecting surface.

Brine for Refrigeration U. S. Patent 1,969,124

A cutectic solution for refrigerating purposes comprises barium chloride 19, potassium chloride 18 and sodium chloride 4 oz. per gallon of water.

Refrigerator Deodorant

Fill a small muslin bag with a good quality of granular activated carbon. The, muslin bag may then be placed in the rear of a lower portion of the ice box and will absorb strong odors which tend to collect.

After six months use, the device may

be reactivated by placing in the oven at 350° F. for about ½ hour.

Increasing Resistance of Magnesium Oxide

U. S. Patent 2.012.897

A process for increasing the electrical resistivity of fused magnesium oxide in an oxidizing atmosphere for approximately 6 hours at a temperature of approximately 2000-2300° F.

Salt Denaturant

Two per cent of wormwood powder is added to salt for industrial use.

Soot Destroyer

Canadian Patent 347,077

Lead Oxide 77 lb. Salt 23 lb.

The above may be diluted with charcoal or sawdust.

Stop Leak Composition

U. S. Patent 1,988,764

A stop leak composition for water circulating systems, comprising as chief ingredients about 4 g. of paper pulp, 5 g. of sifted flax seed, 200 cc, of water, and a small percentage of a preservative.

Temperature Sensitive Compounds

The following color changes induced by temperature changes find applications in many fields:

- 1. Copper Ferroeyanide.- Is mahegany brown at room temperature, becomes brown black on heating, returns to origmal color on cooling.
- Arsenic Bisulphide.—Orange red at room temperatures, changes progressively to dark red and then brown at higher temperatures, returns to original color on cooling.
- Lead Iodide. Original orange changes to dark orange on heating.
- 4. Mercury Subsulphide.—Original yellow changes on heating to orange yellow, then orange, then red.
- Lead Chromate.—Same changes as for mercury subsulphide.
- 6. Tin Subsulphide.—Original brown color (or orange yellow) changes to dark red, then nearly black, on heating. These changes are very temperature sensitive.

- 7. Silver Subiodide.—Green yellow at ordinary temperatures changes to orange when heated.
- 8. Mercury Subiodide.—Original yellowish green changes on heating to orange, red, and brownish red.
- 9. Weak Copper Bromide,—Original lemon-yellow turns to brown when heated, returning to original color when cooled.
- 10. Cobalt Chloride.—Is invisible at ordinary temperatures but becomes blue when heated.
- 11. Mercuric Oxide.—Red at ordinary temperatures, darkens on heating, becomes black eventually.

Thermionic Cathode

U. S. Patent 1,961,122

The filament consists of an alloy of:
Nickel 90 oz.
Iron 7.5 oz.
Titanium 2.5 oz.
Coated with barium oxide.

Protecting Carbide

Carbide will keep indefinitely if sprinkled uniformly with kerosene.

Tooth Desensitizer

(Hartman)

Ether				2	OZ.
Alcohol				1	oz.
Thymol				11/4	oz.
Keep in ered.	a	brown	bottle,	tightly	stop-

Apply inside of tooth by means of a dato of absorbent cotton on a tooth pick. The cavity in which it is applied should be dry to insure lengthy desensitization. Contact should be for 1 to 1½ minutes. The cotton is then removed and the cavity is dried with a blast of hot air.

Denicotinized Cigarettes

Activated charcoal and silica gel is used in individual cigarettes for the absorption of nicotine. Charcoal (0.2 g.) or silica gel (0.1 g.) is an efficient denicotinizer.

Denicotinizing Tobacco U. S. Patent 2,000,855

A method of denicotinizing tobacco comprises the steps of: wetting tobacco containing the usual bacteria, disposing the wetted tobacco loosely in layers and allowing the latter to stand with access of air thereto to produce fermentation of the tobacco, continuously adding acid

to the extent necessary to neutralize the amino bases resulting from the fermentation, and drying the tobacco.

Treating Tobacco for Smoking U. S. Patent 1.972.718

There is added to tobacco about 2% of an alkaline hydrated aluminum silicate which upon the smoking off the tobacco is capable of taking up gases and tarry compounds produced by the combustion.

Water-Softening Compound U. S. Patent 1,952,408

A cake for domestic use, formed by pressure when moist, comprises sodium carbonate 62.5, sodium phosphate 30.0, calcium chloride 5.0, and sodium chloride 2.5%.

Base-Exchange Materials for Water Softening

British Patent 434,663

Raw clay is treated with concentrated hydrochloric or sulphuric acid, the supernatant acid removed, and the clay baked at 550-600° F. for 1 hour. The product is treated with 10% aqueous sodium silicate, then with 2% aqueous sodium aluminate at 100° F., and finally with 5% aqueous sodium chloride to increase the base exchange power.

Water Testing Indicator British Patent 414,866

The dipping rod is coated with a paste made from chalk (16), glycerin (12), a saturated solution of rosin in turpentine (1), and methylene blue dissolved in methylated sprit (1); contact with water lightens the color.

Windshield Anti-Fog Compound Formula No. 1

Windshields may be kept clear of fog, by occasionally wiping them with a cloth prepared by boiling it 10 minutes in a solution of:

Water 5 qt.
Glycerin 1 oz.
Sodium Oleate 1 oz.
Boil together 5 minutes before im-

No. 2
Glycerin 10 oz.
Glycol Boriborate 4 oz.
Sulphonated Castor Oil 10 drops

mersing cloth.

TABLES

Weights and Measures Troy Weight

24 grains = 1 pwt. 20 pwts. = 1 ounce 12 ounces = 1 pound

Apothecaries' Weight

20 grains = 1 scruple 3 scruples = 1 dram 8 drams = 1 ounce 12 ounces = 1 pound

The ounce and pound are the same as in Troy Weight.

Avoirdupois Weight

 $271\frac{1}{32}$ grains = 1 dram 16 drams = 1 ounce 16 ounces = 1 pound 2000 lbs. = 1 short ton 2240 lbs. = 1 long ton

Dry Measure

2 pints = 1 quart 8 quarts = 1 peck 4 pecks = 1 bushel 30 bushels = 1 chaldron

Liquid Measure

4 gills = 1 pint
2 pints = 1 quart
4 quartes = 1 gallon
31½ gals. = 1 barrel
2 barrels = 1 hogshead
1 teaspoonful = ½ oz.
1 tablespoonful = ½ oz.
16 fluid oz. = 1 pint

Circular Measure

60 seconds = 1 minute 60 minutes = 1 degree 360 degrees = 1 circle

Long Measure

12 thches = 1 foot 3 feet = 1 yard 5½ yards = 1 rod 5280 feet = 1 stat. mile 320 rods = 1 stat. mile

Square Measure

144 sq. in. = 1 sq. ft. 9 sq. ft. == 1 sq. yard 30 44 sq. yds. == 1 sq. rod 43,560 sq. ft. == 1 rood 4 roods == 1 rood 4 roods == 1 sq. mile

Metric Equivalents

Length

1 inch == 2.54 centimeters

1 foot == 0.305 meter

1 yard == 0.914 meter

1 mile == 1.609 kilometers

1 centimeter == 0.394 in.

1 meter == 3.281 ft.

1 kilometer == 0.621 mile

Capacity

1 U. S. fluid oz. == 29.573 milhiliters
1 U. S. hquid qt. == 0.946 liter
1 U. S. dry qt. == 1.101 liters
1 U. S. gallon == 3.785 liters
1 U. S. bushel == 0.3524 heetoliter
1 u. m. == 16.4 cu. centimeters
1 milhiliter == 0.034 U. S. fluid ounco
1 liter == 1.057 U. S. hquid qt.
1 liter == 0.908 U. S. dry qt.
1 liter == 0.264 U. S. gallon
1 heetoliter == 2.838 U. S. bu.
1 cu. centimeter == .001 cu. m.
1 liter == 1.000 milliliters or 100 cu. c.
1 liter == 1.000 milliliters or 100 cu. c.

Weight

1 grain = 0.065 gram
1 apoth, seruple = 1.206 grams
1 av. oz. = 28.356 grams
1 troy oz. = 31.103 grams
1 av. lb. = 0.454 kilogram
1 troy lb. = 0.373 kilogram
1 gram = 15.432 grains
1 gram = 0.772 apoth, seruple
1 gram = 0.035 av. oz.
1 gram = 0.032 troy oz.
1 kilogram = 2.205 av. lbs.
1 kilogram = 2.205 av. lbs.

386 THE CHI	EMICAL	FORMULARY	
Approximate pH Values		Beets	4.9-5.5
		Blackberries	3.2-3.6
The following tables give appro	ximate	Bread,*white	5.0-6.0
pH values for a number of sub-	stances	Butter	6.1-6.4
such as acids, bases, foods, bio	logical	Cabbage	5.2-5.4
fluids, etc. All values are rounded		Carrots	4.9-5.3
the nearest tenth and are based on	meas-	Cheese	4.8-6.4
urements made at 25° C.	- 1	Cherries	
pH Values of Acids	1		3.2-4.0
•		Cider	2.9-3.3
Hydrochloric, N	. 0.1	Corn	6.0-6.5
Hydrochloric, 0.1N	. 1.1	Crackers	6.5-8.5
Hydrochloric, 0.01N	. 2.0	Dates	6.2-6.4
5011D011F1C. IV		Eggs, fresh white	7.6-8.0
Sulphuric, 0.1N	1.2	Flour, wheat	5.5-6.5
Sulphuric, 0.1N	. 2.1	Gooseberries	2.8-3.0
Orthophosphoric, 0.1N	1.5	Grapefruit	3.0-3.
Sulphurous, 0.1N		Grapes	3.5-4.5
Oxalic, 0.1N	. 1.6	Hominy (lye)	6.8 - 8.0
Tartaric, 0.1N	2.2	Jams, fruit	3.5-4.0
Malic. 0.1N		Jellies, fruit	2.8-3.
Malic, 0.1N Citric, 0.1N Formic, 0.1N	2.2	Lemons	2.2-2.
Formie 0.1N	2.3	Limes	1.8-2.
Lactic, 0.1N	2.4	Maple Syrup	6.5-7.
Agotic N	2.4	Milk, cows	6.3-6.
Agotio 01N	2.9	Olives	3.6-3.
Agotio 0.01N	3.4	Oranges	3.0 - 4.
Acetic, N Acetic, 0.1N Acetic, 0.01N Benzoic, 0.1N	3.1	Oysters	6.1 - 6.
Alum 0 1N	3.2	Peaches	3.4-3.
Alum, 0.1N		Pears	3.6-4
Carbonic (saturated)	4.1	Pens	5.8-6.
Araniana (anturated)	5.0	Pickles, dill	3.2 - 3.
Arsenious (saturated)		Pickles, sour	3.0-3.
Hydrocyanic, 0.1N		Pimento	4,6-5.
Boric, 0.1N	0.5	Plums	2.8-3.
pH Values of Bases		Potatoes	5.6-6.
•		Pumpkin	4.8 - 5
Sodium Hydroxide, N	. 14.0	Raspherries	3.2 - 3.
Sodium Hydroxide, 0.1N	. 13.0	Rhubarb	3.1-3
Sodium Hydroxide, 0.01N		Salmon	6.1-6.
Potassium Hydroxide, N	. 14.0	Sauerkraut	3.4-3.
Potassium Hydroxide, 0.1N	. 13.0	Shrimp	6.8-7.
Potassium Hydroxide, 0.01N	. 12.0	Soft Drinks	2.0-4
Lime (saturated)	. 12.4	Spinach	5.1-5
Sodium Metasilicate, 0.1N	. 12.6	Squash	5.0-5
Trisodium Phosphate, 0.1N	. 12.0	Strawberries	3.0 - 3
Sodium Carbonate, 0.1N	. 11.6	Sweet Potatoes	5.3-5
Ammonia, N	. 11.6	Tomatoes	4.0-4
Ammonia, 0.1N	. 11.1	Tuna	5.9-6
Ammonia, 0.1N	. 10.6	Turnips	5.2-5
Potassium Cyanide, 0.1N	. 11.0	Vinegar	2.4-3
Magnesia (saturated)		Water, drinking	6.5-8
Sodium Sesquicarbonate, 0.1N	. 10.1	Wines	2.8-3
Ferrous Hydroxide (saturated) .	. 9.5		
Calcium Carbonate (saturated)		pH Values of Biologic Mat	erials
Borax, 0.1N	. 9.2	Blood, plasma, human	7.3-7
Sodium Bicarbonate, 0.1N		Spinal Fluid, human	7.3-7
			6.9-7
pH Values of Foods		Blood, whole, dog	6.5-7
	2.9-3.3		1.0-3
Apples		Gastric Contents, human	
Apricots	3.6-4.0	Duodenal Contents, human	4.8-8 4.6-8
Asparagus	5.4-5.8	Feces, human	4.8-8
Bananas	4.5-4.7 5.0-6.0	Urine, human Milk, human	6.6-7
Beans	4.0-5.0	Bile, human	6.8-7
Beers	3.0-0.0	I Due, namen	0.0-1

CONVERSION OF THERMOMETER READINGS

F°	C _o	F°	C.	F°	C.	F°	C.	F°	C.	F°	C.	
-40 -38 -36 -34 -32	-40.00 -38.89 -37.78 -36.67 -35.56	30 31 32 33 34	-1 11 -0.56 0.00 0.56 1.11	80 81 82 83 84	26.67 27.22 27.78 28.33 28.89	250 255 260 265 270	121.11 123.89 126.67 129.44 132.22	500 505 510 515 520	260 00 262.78 265.56 268.33 271.11	900 910 920 930 940	487.78 493 33 498.89	
$ \begin{array}{r} -30 \\ -28 \\ -26 \\ -24 \\ -22 \end{array} $	-34.44 -33.33 -32.22 -31.11 -30.00	35 36 37 38 39	1.67 2.22 2.78 3.33 3.89	85 86 87 88 89	29.44 30.00 30.56 31.11 31.67	275 280 285 290 295	135.00 137.78 140 55 143.33 146.11	535	273.89 276.67 279.44 282.22 285.00	950 960 970 980 990	510.00 515.56 521.11 526.67 532.22	
-20 -18 -16 -14 -12	-28.89 -27.78 -26.67 -25.56 -24.44	40 41 42 43 44	4.44 5.00 5.56 6.11 6.67	90 91 92 93 94	32.22 32.78 33.33 33.89 39.44	300 305 310 315 320	148.89 151.67 154.44 157.22 160.00	555 560 565	287.78 290.55 293.3 296.1 298.89	1050 1100	537.78 565.56 593.33 621.11 648.89	
-10 - 8 - 6 - 4 - 2	-23.33 -22.22 -21.11 -20.00 -18.89	45 46 47 48 49	7.22 7.78 8.33 8.89 9.44	95 96 97 98 99	35.00 35.56 36.11 36.67 37.22	330 335	162.78 165.56 168.33 171.11 173.89	580 585 590	301.67 304.44 307.22 310.00 312.78	1300 1350 1400	676.67 704.44 732.22 760.00 787.78	
0 1 2 3 4	-17.78 -17.22 -16.67 -16.11 -15.56	50 51 52 53 54	10.00 10.56 11.11 11.67 12.22	100 105 110 115 120	37.78 40.55 43.33 46.11 48.89	350 355 360 365 370	176.67 179.44 182.22 185.00 187.78	600 610 620 630 640	315.56 321.11 326.67 332.22 337.78	1550 1600 1650	871.11 898.89	
5 6 7 8 9	-15.00 -14.44 -13.89 -13.33 -12.78	55 56 57 58 59	12.78 13.33 13.89 14.44 15.00	125 130 135 140 145	51.67 54.41 57.22 60.00 62.78	375 380 385 390 395	190.55 193.33 196.11 198.89 201.67	670	343.33 348.89 354.44 360.00 365.56	1800 1850 1900	954.44 982.22 1010.00 1037.78 1065.56	
10 11 12 13 14	-12.22 -11.67 -11.11 -10.56 -10.00	60 61 62 63 64	15.56 16.11 16.67 17.22 17.78	150 155 160 165 170	65.56 68.33 71.11 73.89 76.67	400 405 410 415 420	204.44 207.22 210.00 212.78 215.56	700 710 720 730 740	371.11 376.67 382.22 387.78 393.33	2050 2100 2150	1093.33 1121.11 1148.89 1176.67 1204.44	
15 16 17 18 19	- 9.44 - 8.89 - 8.33 - 7.78 - 7.22	65 66 67 68 69	18.33 18.89 19.44 20.00 20.56	175 180 185 190 195	79.44 82.22 85.00 87.78 90.55	425 430 435 440 445	218.33 221 11 223.89 226.67 229.44	750 760 770 780 790	308.89 404.44 410.00 415.56 421.11	2300 2350 2400	1232.22 1260.00 1287.78 1315.56 1343.33	
20 21 22 23 24	- 6.67 - 6.11 - 5.56 - 5.00 - 4.44	70 71 72 73 74	21.11 21.67 22.22 22.78 23.33	200 205 210 215 220	93.33 96.11 98.89 101.67 104.44	450 455 460 465 470	232 22 235 00 237.78 240 55 243.33	800 810 820 830 840	426 67 432 22 437.78 443.33 448.89	2550 2600 2650	1371.11 1398.89 1426 67 1454.44 1482.22	
25 26 27 28 29	- 3.89 - 3.33 - 2.78 - 2.22 - 1.67	75 76 77 78 79	23.89 24.44 25.00 25.56 26.11	225 230 235 240 245	107.22 110.00 112.78 115.56 118.33	475 480 485 490 495	246 11 248.89 251.67 254.44 257.22	850 860 870 880 890	454.44 460.00 465.56 471.11 476.67	2800 2850 2900	1510.00 1537.78 1565.56 1593.33 1621.11	

ALCOHOL PROOF AND PERCENTAGE TABLE

	ALCOHOL	I I WOOF AND	I DIVIDINIA	IL IADLE	
U.S. Proof at 60° F.	Per cent Alcohol by Volume at 60° F.	Per cent Alcohol by Weight	U.S. Proof at 60° F.	Per cent Alcohol by Volume at 60° F.	Per cent Alcohol by Weight
0	0.0	0.00	58	29.0	23.82
i	0.5		59	29.5	20.02
2	1.0	0.80	60	30.0	24.67
2 3 4	1.5		61	30.5	
4	2.0	1.59	62	31.0	25.52
5	2.5		63	31.5	
6	3.0	2.39	64	32.0	26.38
7 8	3.5		65	32.5	
9	4.0 4.5	3.19	66	33.0	27.24
10	5.0	4.00	67 68	33.5 × 34.0	28.10
iĭ	5.5	4.00	69	34.5	28.10
12	6.0	4.80	70	35.0	28.97
13	6.5		71	35.5	
14	7.0	5.61	72	36.0	29.84
15	7.5		73	36.5	
16	8.0	6.42	74	37.0	30.72
17 18	8.5	7 .00	75	37.5	
19	9.0 9.5	7.23	76	38.0	31.60
20	10.0	8.05	77 78	38.5 39.0	20.40
21	10.5	0.00	79	39.5	32.48
22	11.0	8.86	80	40.0	33.36
23	11.5		81	40.5	
24	12.0	9.68	82	41.0	34.25
25	12.5		83	41.5	
26	13.0	10.50	84	42.0	35.15
27	13.5	11.00	85	42.5	
28 29	14.0 14.5	11.32	86 87	43.0	36.05
30	15.0	12.14	88	43.5 44.0	36.96
31	15.5	12/11	89	44.5	30.90
32	16.0	12.96	90	45.0	37.86
33	16.5		91	45.5	
34	17.0	13.79	92	46.0	38.78
35	17.5		93	46.5	***************************************
36 37	18.0	14.61	94	47.0	39.70
38	18.5 19.0	15.44	95 96	47.5	10.00
39	19.5	10.44	97	48.0 48.5	40.62
40	20.0	16.27	98	49.0	41.55
41	20.5		99	49.5	
42	21.0	17.10	100	50.0	42.49
43	21.5		101	50.5	
44	22.0	17.93	102	51.0	43.43
45 46	22.5 23.0	18.77	103	51.5	
47	23.5	10.77	104 105	52.0 52.5	44.37
48	24.0	19.60	106	53.0	45.33
49	24.5		107	53.5	10.00
50	25.0	20.44	108	54.0	46.28
51	25.5		109	54.5	
52	26.0	21.28	110	55.0	47.24
53 54	26.5	00.12	111	5 5.5	
55	27.0 27.5	22.13	112	56.0	48.21
56	28.0	22.97	113 114	56.5 57.0	10.10
57	28.5		115	57.5	49.19
				00	

		1111	BLES		
U. S. Proof	Per cent Alcohol by Volume	Per cent Alcohol	U. S. Proof	Per cent Alcohol by Volume	Per cent Alcohol
at 60° F.	at 60° F.	by Weight	at 60° F.	at 60° F.	by Weight
116	58.0	50.17	159	79.5	
117	58.5		160	80.0	73.53
118	59.0	51.15	161	80.5	
119	59.5		162	81.0	74.69
120	60.0	52.15	163	81.5	
121	60.5		164	82.0	75.86
122	61.0	53.15	165	82.5	
123	61.5		166	83.0	77.04
124	62.0	54.15	167	83.5	
125	62.5		168	84.0	78.23
126	63.0	55.16	169	84.5	-
127	63.5		170	85.0	79.44
128	64.0	56.18	171	85.5	*****
129	64.5		172	86.0	80.62
130	65.0	57.21	173	86.5	
131	65.5		174	87.0	81.90
132	66.0	58.24	175	87.5	
133	66 .5		176	88.0	83.14
134	67.0	59.28	177	88.5	
135	67.5		178	89.0	84.41
136	68.0	60.32	179	89.5	
137	68.5		180	90.0	85.69
138	69.0	61.38	181	90.5	
139	6 9.5		182	91.0	86.99
140	70.0	62.44	183	91.5	
141	70.5		184	92.0	88.31
142	71.0	63.51	185	92.5	
143	71.5		186	93.0	89.65
144	72.0	64.59	187	93.5	
145	72.5		188	94.0	91.02
146	73.0	65.67	189	94.5	
147	73.5		190	95.0	92.42
148	74.0	66.77	191	95. 5	
149	74.5		192	96.0	93.85
150	75.0	67.87	193	96.5	
151	75.5	-	194	97.0	95.32
152	76.0	68.92	195	97.5	
153	76.5	-	196	98.0	96.82
154	77.0	70.10	197	98.5	
155	77.5		198	99.0	98.38
156	78.0	71.23	199	99.5	
157	78.5		200	100.0	100.00
158	79.0	72.38			

The following table gives some com-
mon buffer systems and the approximate
pH of maximum buffer capacity. The
zone of effective buffer action will vary
with concentration but the general aver-
age will be ± 1.0 pH from the value
given, for concentrations approximately
0.1 molar.

chloric Acid	2.8
Primary Potassium Citrate	3.7
Acetic Acid-Sodium Acetate	4.6

Potassium Acid Phthalate-Sodium	
Hydroxide	5.0
Secondary Bodium Citrate	5.0
Carbonic Acid-Bicarbonate	6.5
Primary Phosphate-Secondary Phos-	•••
phate	6.8
Primary Phosphate Sodium Hydrox-	
ide	6.8
Borie Acid-Borax	8.5
Borax	9.2
Boric Acid Sodium Hydroxide	9.2
Bicarbonate-Carbonate	10.2
Secondary Phosphate-Sodium Hy-	
droxide	11.5
Courtery of W. A. Taylor & Com	0674

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Amer. Dyestuff Reporter
Amer. Electrop. Society
Amer. Paint Jol.
Amer. Perfumer
Amer. Photography
Amer. Wool & Cotton Reporter
Anal Fis. Quim.

Ault & Wiborg Varnish Wks. Handbook

Baker's Helper
Bakers Review
Baker's Weekly
Better Enameling
Bottler & Packer
Boyce Thompson Inst.
Browers' Tech. Review
Brick & Clay Record
Br. Jol. Dent. Science
Brit. Jol. of Photography
Brit. Medical Jol.
Bull. Imp. Hyg. Lab.
Bulletin of Imperial Instituto
Bull. So. Franc. Phot.

Camera
Camera (Luzern)
Canner
Cement & Cement Mfr.
Chemical Abstracts
Chemical Analyst
Chemical Industries
Chemical Weekblad
Chem. Zent.
Chemist & Druggist
Combustion
Confectioner's Jol.
Cramer's Manual

Dairy World
Danek, Tide, Farm
Dental Lab'y Review
Devt. Part. Zeitung
Drug & Cosmetic Industry
Druggists Circular
Drugs, olls, & Paints

Eastman Kodak Co. Electric Journal

Farbe v. Lacke

Farben Zeitung
Farming S. Africa
Fein Mechanic v. Prazision
Fettchem, Umschan
Fils & Tissus
Focus
Food Manufacture
Fruit Products Jol.

Gelatin, Leim, Klebstoffe Glass Industry

Hawaiian Planters' Record

Ice Cream Review
India Rubber World
Indian Lac Research Inst.
Industrial Chemist
Industrial Finishing
Int'l Tin Res. & Dev. Council

Jol. Amer. Dental Assn.
Jol. Amer. Medical Assn.
Jol. Chinese Chem. Soc.
Jol. Federation Curriers
Jol. Federation Light Leather Tanners
Jol. Ind. & Eng. Chemistry
J. Res. Nat. Bur. Standards
Jol. Rubber Industry
J. Russ. Rubber Ind.
Jol. Soc. Leather Trades
Jol. Soc. Rubber Ind. Japan

Keram Steklo Khimstroi Kozhevenna-Obuvnaya Prom. Kunstdunger, Und Leim

Lakokras, Ind. Leather Trades Review Les Mat. Grasses Lithographic Tech. Foundation

Malayan Agric. Jol.
Manufacturing Chemist
Meat
Meat Merchandising
Melliand
Metal Industry
Metall und Ers
Metallurg
Metallurgist
Metals & Alloys

Mich. Agric. Exp. Sta. Monatschr. Textil-Ind. Munic. Eng. San. Record

Nat'l Butter & Cheese Jol. Nat'l Provisioner Nickelsworth Nitrocellulose

Ober Flachen Tach. Oil & Color Trades Jol. Oil & Soap

Paper Trade Jol. Parfum Mod. Penture, Pigments, Vernis Phar. Acta Helva Pharmaceutical Jol. Phot. Abstracts l'hot, Ind. Phot. Korr. Photog. Kronik Phot. Rev. Photo Rundschau l'hysics Phytopathology Plater's Guide Book Portland Cement Assn. Power Practical Druggist

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Textile Colorist Textile Mfr. Textile Recorder

U. S. Department of Agriculture U. S. Bureau of Mines U. S. Bureau of Standards

Veneers and Plywood

Z. Elektrochem. Zeit. Unters. Lebensm.

COMMON NAMES OF CHEMICAL PRODUCTS

.

Capticum Red Pepper Carbolic Acid Phenol Carragheen Irish Moss		
CatechuCutch		
Caustic Potash Potassium Hydrovide		
Caustic SodaSodium Hydroxide		
Ceresin WaxOzokerite and Paraffin Mixture ChalkCalcium Carbonate		
China Clay		
China Wood OilTung Oil		
Chinese Wax		
Cholestrin		
Chrome Green		
Cinnabar Mercuric Sulphide		
Citronella Oil		
Colloidal ClayBentonite		
Collodion		
Cologne Spirits Ethyl Alcohol (pure)		
Colophony Rosin Pine Resin		
Columbian Spirits		
Colza Oil		
Copper Aceto Arsenite		
Copper Arsenite		
Corn SugarDextrose Corn SyrupGlucose		
Corrosive Sublimate		
Corundum		
Cresol		
Crude OilPetroleum (crude)		
CyanamidCalcium Cyanamide		
D		
D		
Dead OilCreosote Oil		
Dead Oil		
Dead Oil		
Dead Oil		
Dead Oil		
Dead Oil		
Dead Oil		
Dead Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Decad Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		
Dead Oil Creosote Oil		

G

Galena Lead Sulphide Glance Pitch Manjak Glass, Water Sodium Silicate Glauber's Salt Sodium Sulphate Glycerin Glycerol Glycol Ethylene Glycol Graphite Plumbago Green Soap Soft Soap Green Old Ferrous Sulphate Ground Nut Oil (Arachi's Oil) Peanut Oil Gum Lac Shellac Gun Cotton Nitro-Cellulose Gypsum Calcium Sulphate
н
Heavy Spar
1
Ichthyol
К
Kauri Gum Copal, Gum Kieselguhr { Tripoli Diatomaceous Earth
L
Lanum Lanolin Lead Chromate Chrome Yellow Lead Sulfate, Basic Whitelead, Sublimed Lemon, Salts of Potassium Binoxalate Licorice Glycyrrhiza Ligroin, Light Petroleum Ether Lime Calcium Oxide Lime, Slaked Calcium Gride Limestone Calcium Carbonate Litharge Lead Monoxide Liver of Sulphur Potassium Sulphide Lunar Caustic Silver Nitrate Lye Sodium Hydroxide
M
Magnesium, Calcined Magnesium Oxide Magnesium Silicate Talcum Maize Oil Corn Oil

Milk Sugar Lactose Mineral Pitch Asphalt Minium Lead Oxide (red) Mirbane Oil Nitrobenzol Muriatic Acid Hydrochloric Acid Myrtle Wax Bayberry Wax
N
Naphtha, Solvent Coal Tar Naphtha Naples Yellow Lead Antimonnte Nickel Salta, Double Nickel Ammonium Sulphate Nickel Salta, Single Nickel Sulphate Niter Potassium Nitrate Niter Cake Sodium Bisulphate Nitrocellulose (soluble cotton) Pyroxylin
•
Oleic Acid Red Oil Jlein Glyceryl Tri-oleate (natural) Jleum Sulphuric Acid (fumng) Jlive Oil Sweet Oil Jrange Mineral Orange Red Lead Oxide Jrpiment Arsenous Sulphide (yellow)
P
Paraffin Oil Mineral Oil
Paraffin Oil Sineral Oil Petrolatum, Liquid Paris White Whiting Petrolatum, Liquid Petrolatum, Liquid Petrolatum, Liquid Petrolatum, Liquid Petrolatum Carbonato Petrole Gasoline Petroletum Petroleum Jelly Paster of Paris Calcium Sulphate plus 1 mol. water Potassium Bicarbonate Salaterus Prussian Blue Ferric Ferrocyanido Prussiate of Potash, Red Potassium Ferricyanido Prussiate of Potash, Yellow Potassium Ferrocyanido Prussiate Acid Hydrocyanic Acid Pyroligneous Acid Wood Vinegar
Q
uicklimeCalcium Oxide uickailverMercury
R
ed Oxide
8
accharine

Saltpeter Scale Wax Silica Sod Oil Soda Ash Sodium Bisulphite Sodium Phosphate, Dibasic Sodium Phosphate, Monobasic Sodium Phosphate, Tribasic Sodium Phosphate, Tribasic Sodium Thiosphate Sperm Oil Spirits of Turpentine Stannous Chloride Stearin Storax Sucrose Sugar of Lead Sulfonated Castor Oil Sulphur Olive Oil Sulphuric Acid Sulphuric Ether	.Paraffin Wax (low melting) .Silicon Dioxide .Degras .Sodium Carbonate, Anhydrous .Sodium Acid Sulphate .Disodium Phosphate .Disodium Phosphate .Trisodium Phosphate .Trisodium Phosphate .Hypo .Whale Oil .Turpentine .Tin Crystals .Tristearin .Styrax Cane Sugar Leed Acetate .Turkey Red Oil .Olive Oil Foots
	-
	T
marm.	
TNT	. Trinitrotoluene
Tartar Emetic	Antimony Potassium Tartrate
I etrain	Totachardae Menhahalas
Theobroma Oil	Coses Dutter
Titanium Dioxide	Oacao Dutter
Tolune	Titanium Oxide
Toluene	Toluol
Triacetin	Glycerol Triacetate
Trinitrophenol	Pierie Acid
•	
	V
Verdigris	Copper Acetate Basic
Verdigris	Mercuric Sulphide Red
	Surpinde, 200
	W
TITL-1. 011	
Whale Oil	Train Oil
White Arsenic	Arsenic Trioxide
White Bole	Vaslia
White Lead	Lead Carbonata Rosia
White Metal	Rabbitt Motal
White Wax	Posses (blooks)
Whiting	Challe Defend
Wintergreen Oil Symthetic	Onnik, Renned
Wintergreen Oil, Synthetic	Metnyl Balicylate
Wood Alcohol	Methyl Alcohol
	Y
	· ·
Vacca Cum	
INCER CHIII	Accrecides Com
Yacca Gum	Accroides Gum
iacca dum	Accroides Gum
iacca Gum	
	z
Zinc White	Z Zine Ovida
Zinc White	Z Zine Ovida
	Zinc Oxide

TRADE NAMED CHEMICALS

During the past few years, the practice of marketing raw materials, under names which in themselves are not descriptive chemically of the products they represent, has become very prevalent. No modern book of formulae could justify its claims either to completeness or modernity without numerous formulae containing these so-called "Trade Names."

Without wishing to enter into any discussion regarding the justification of "Trade Names," the Editors recognize the tremendous service rendered to commercial chemistry by manufacturers of "Trade Name" products, both in the physical data supplied and the formulation suggested.

Deprived of the protection afforded their products by this system of nomenclature, these manufacturers would have been forced to stand helplessly by while the fruits of their labor were being filched from them by competitors who, unhampered by expenses of research, experimentation and promotion, would be able to produce something "just as good" at prices far below those of the original producers.

That these competitive products were "just as good" solely in the minds of the imitators would only be evidenced in costly experimental work on the part of the purchaser and, in the meantime irreparable damage would have been done, to the truly ethical product. It is obvious, of course, that under these circumstances, there would be no incentive for manufacturers to develop new materials.

Because of this, and also because the "Chemical Formulary" is primarily concerned with the physical results of compounding rather than with the chemistry involved, the Editors felt that the inclusion of formulae containing various trade name products would be of definite value to the producer of finished chemical materials. If they had been left out many ideas and processes would have been automatically eliminated.

As a further service a list of the better known "trade name" products is appended together with the suppliers of these materials. The number after each trade name refers to the supplier given below with the corresponding number.

TRADE NAMES

Α	Ascarite
	Astrulan 6
A Syrup	Atrapol
Abalyn 79	Aurosal
Abopon 70	Avonac
Accelerator 808 51	
Accelerator 833 51	В
Acetoin 94	Badex
Acidolene 47	Bakelite
Acto149	Bardol
Adheso Wax 70	Barretan 16
A.D.M. No. 100 Oil 10	Beckacite
Aerogel101	Beckolin
Agerite Powder163	Beckosol
Akcocene 6	Bensapol
Alba-Floc	Beutene
Albasol106	Blandol
Albatex 38	Blendene 70
Albertol	Bludtan
Albinol	Bordow 49
Albolith	Borol
Albone "C" 51	Bromo "Acid"
Albusol 96	Brosco
Aldehol 87	Butalyde42
Aldol181	Butyl Carbitol
Alkanol 51	Butyl Cellosolve
Alloxan 20	•
Aloxite 29	C
Alphasol 6	Cadalyte 73
Altax163	
	Cadmonth
Alugel104	Cadmolith
Alugel	Calcoloid
Alugel .104 Amberette .154 Amberol .125	Calcoloid 25 Calcene 41
Alugel 104 Amberette 154 Amberol 125 Ambreno 51	Calcoloid
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amo Acetate 88	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165
Alugel 104 Amberette 154 Amborol 125 Ambreno 51 Amo Acetate 88 Amandol 51 Amidine 26	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amoo Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Capticol 28 Carboxide 28 Caseo 30 Catalpo 102 CCH 98
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amoo Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 8	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captico 28 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Capticol 28 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amcon 51 Amcon 61 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censteric 32 Cerciose 44 Cereps 170
Alugel 104 Amberol 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55 Aquasol 6	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captice 185 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censteric 32 Cerelose 44
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquaspel 114 Aquasoe 55 Aquasol 6 Arapali 129	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captice 155 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censteric 32 Cerelose 44 Cereps 170 Ceresalt 53 Chlorex 28
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Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55 Aquasol 6 Arapali 129 Araskleen 101 Archer-Daniels No. 635 10	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censteric 32 Cerelose 44 Cereps 170 Ceresalt 53 Chlorax 28 Chlorasol 28 Chremitz White 56
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Ambreno 51 Amcon 51 Amea Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55 Aquasol 6 Aquasol 6 Arapali 129 Araskleen 101 Archer-Daniels No. 635 10 Archer-Daniels No. 635 10 Archer-Daniels No. 635 10	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censele 44 Cereps 170 Ceresalt 53 Chlorex 28 Chlorasol 28 Chremitz White 56 Cinchophen 25
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amcon 51 Amcon 51 Amidine 26 Anchoracel 27 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquanel 70 Aquapel 114 Aquarome 55 Aquasol 6 Arapali 129 Araskleen 101 Archer-Daniels No. 635 10 Ardex 51 Ardes 114 Ardex 51	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censteric 32 Cerese 44 Cereps 170 Ceresalt 53 Chlorasol 28 Chromatiz White 56 Cinchophen 25 Coblac 19
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55 Aquamel 129 Araskleen 101 Archer-Daniels No. 635 10 Aridex 51 Arochlor 153	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celoscour 3 Celite 85 Cellosolve 28 Censeric 32 Cerelose 44 Cereps 170 Ceresalt 53 Chlorex 28 Chromable 28 Chromolit 48 Coblae 19 Cominol 43
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amcon 51 Amcon 51 Amcon 51 Amidine 26 Amchoracel 27 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55 Aquasol 6 Arapali 129 Araakleen 101 Archer-Daniels No. 635 10 Archer-Daniels-Midland Oil 10 Aridex 51 Arcolor 153 Arcolor 153 Arcolor 153 Arcolor 163	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celascour 3 Celite 85 Cellosolve 28 Censteric 32 Cerelose 44 Cereps 170 Ceresalt 53 Chlorasol 28 Chremnitz White 56 Cinchophen 25 Coblae 19 Cominol 43 Coppercide 83
Alugel 104 Amberette 154 Amberol 125 Ambreno 51 Amco Acetate 88 Amandol 51 Amidine 26 Anchoracel 2p 7 Anhydrone 14 Ansol 161 Antidolorin 58 Apcothinner 8 Aqualoid 86 Aquamel 70 Aquapel 114 Aquarome 55 Aquamel 129 Araskleen 101 Archer-Daniels No. 635 10 Aridex 51 Arochlor 153	Calcoloid 25 Calcene 41 Calgon 22 Calorite 148 Captax 165 Carbitol 28 Carboxide 28 Casco 30 Catalpo 102 CCH 98 Celoscour 3 Celite 85 Cellosolve 28 Censeric 32 Cerelose 44 Cereps 170 Ceresalt 53 Chlorex 28 Chromable 28 Chromolit 48 Coblae 19 Cominol 43

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Cromodine	ì
Cryptone	Idalol
Cumar	Igepon
Cyclamal	IG Wax O 65
Cycline	Indian Red 19
Cymanol 82	Indigusols
D	Indur
•	Isolene
Darco 45	2501020
Diamond K Linseed Oil145	J
Dionin100	
Discolite	Jasmogene165
Disperso	к
Distoline	^
Duolith 90	Kalite
Duphax146	Karo 44
Duphonol 51	Kellogg Kuo145
DuPont Rubber Red 51	Kellogg Varnish Oil145
Durez 68	Kerol 21
	Kilfonm 4
E	Kolineum
	Kopol
Fastman Products	Koreon103
Elame 54	Kryocide118
Erio Chrome Dyes	
Esterol115	L
Estersol161	Lactol Spirita 85
Ethox184	
Ethyl Parasept	
Ethyl Protol	Lanctte Wax
Eulan 65	Le Page's Cement132
F	Leukonin
•	Lewisol
Factolac 81	Lindol
Falba Absorption Base	Lohrinol
Feectol131	Lucidol94
Fer-ox173	Lysol
Ferrox150	17,.01
Fixalt101	M
Flexoresin 70	30
Fyrex166	Mapico
	Mellittis 69
G	Merpentine
Janaina) E1	Methyl Cellosolve
Gardinol	Metso
Gastex	
Gilsonite	Monex108
	N
	Naccon105
Glyptal 66	Naccolene
Glutrin128 Guai-a-phene	National Oil Red
Guantal	Nekal 65
neriter	Nelgin 70
н	Neomerpin 51
	Neutroleum 60
Halowax 76	Nevindene109
Hercusol 79	Nevinol109
Hydralite C 65	Nipagen 71
Hydristear172	Nitramon 51
Hydromalin 70	Nu-char 82
Hydroresin	Nulomoline111
Hydrowax 70	Nuodex112

100 IRADE	IVAMIA
O Oildag 1	"S" Syrup
Oil Root Beer C	Stearol
Olate	Stripolite
Ondulum 70	Stripper T. S
Opal Wax 51	Sulfo Turk C
Osmo-Kaolin 57	Sulphoricinol
Oxynone131	Sunoco Spirits152
_	Burfex183
P	Syntex
Parachol 70	•
Paracide	T
Para-dor	
Para-flux	Tanax 6
Paramet	Teglac 6
Paranol115	Telloy
Paris Black 19	Thionex
Paris White144	Timonex
Paroil4	Titanox
Peerless Clay	Ti-Tone 90
Pentrol	Tonsil
Perchloron	Tornesit 79
Perrol	Triclene 51
Petrohol147	Tuads
Pharmasol	Tunguran A 2
Plastogen	Turkelene 70
Plioform 72	11
Pliolite 72	•
Proofit 70	Ultrasene 12
Proxate 93	Unilith159
Puerine	Ureka C
Pyrax	Ursulin 6
Pyrefume	Uversol 2
Pyrethrol	v
•	· · · · · · · · · · · · · · · · · · ·
Q	Valex
Quakersol	Vandex 163 Vanillal 142
•	Varcum
R	Varnolene
Rapidase	Vaseline
Rausene	Vaso167
Resin R-H-35	Vatsol
Resinox126	Vinapas 2
Revertex127	Vinsol 78
Rezyl 6	Vinylite 28
Rodo163	Volclay
Roseol 95	Varioz100
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O-L-II- Ollin-	Wyo-Jel
Schultz Silica 34 Sellatan A 61	1,000 01
Serinol	x
Sherpetco141	w1 ~
Silex	Xerol
Soligen 2	X-13 6
Solosone 51	z
Bolwax124	
Speron 24	Zimate16
SRA Black 3	Zopaque17

SUPPLIERS OF TRADE NAME CHEMICALS

1. Acheson Graphite Corp., Niagara Falls, N. Y.
2. Advance Solvents & Chem. Corp., New York City
3. American Aniline Products, Inc., New York City
4. American Chem. Prod. Co., Rochester, N. Y.
5. American Colloid Co., Chicago, Ill.
6. American Cyanamid & Chem. Co., New York City
7. Anchor Chem. Co., Manchester, England
8. Anderson Prichard Oil Corp., Oklahoma City, Okla.
9. Anabez-Rigiel Corp., Rosebank, N. Y.

Anabez-Rigiel Corp., Rosebank, N. Y. 9. Ansbacher-Siegle Corp., Rosebank, N. Y.
10. Archer-Daniels-Midland Co., Minneapolis, Minn.
11. Arkanssa Co., New York City 11. Arkansas Co., New York City
12. Atlantic Refining Co., Phila., Pa.
13. Bakelite Corp., New York City
14. Baker, J. T. Chem. Co., Phillipsburg, N. J.
15. Barber Asphalt Co., Phila., Pa.
16. Barrett Co., New York City
17. Beek Kolles & Co. 14. Baker, J. T. Chem. Co., Phillipaburg, N. J.
15. Barber Asphalt Co., Phila., Pa.
16. Barrett Co., New York City
17. Beck, Koller & Co., Detroit, Mich.
18. Bick & Co., Inc., Reading, Pa.
19. Binney & Smith, New York City
20. British Drug Houses, Ltd., London, England
21. Bud Aromatic Chem. Co., Inc., New York City
22. Buromin Corp., Pittsburgh, Pa.
23. Bush, W. J. & Co. Inc., New York City
24. Cabot, Godfrey L. Inc., Boston, Mass
25. Calco Chem. Co., Bound Brook, N. J.
26. Campbell, John & Co., New York City
27. Carbic Color & Chem. Co., New York City
28. Carbide & Carbon Chem. Corp., New York City
29. Carborundum Co., Niagara Falls, N. Y.
30. Casein Mfg. Co., New York City
31. Cellulod Corp., Newark, N. J.
32. Century Stearie Acid & Candle Wks., New York City
33. Champion Fibre Co., Canton, No. Car.
34. Chaplin-Bibbo, New York City
35. Chemical & Pigment Co., Inc., Scranton, Pa.
36. Chemical Solvents Inc., New York City
37. Chesebrough Mfg. Co., New York City
38. Ciba Co., Inc., New York City
39. Colgate-Palmolive-Peet Co., Jersey City, N. J.
40. Colledge, E. W., Inc., Cleveland, O.
41. Columbia Alkali Corp., New York City
42. Commercial Solvents Corp., Terre Haute, Ind.
43. Commonwealth Color & Chem. Co., Brooklyn, N. Y.
44. Corn Products Refining Co., New York City
45. Darco Sales Corp., New York City
46. Deep Rock Oil Corp., Chicago, Ill.
47. Dennis, Martin & Co., New York City
48. Dodge & Olcott Co., New York City
49. Dow Chem. Co., Midland, Mich.
40. Ducas, B. P. Co., New York City
41. Dennis, Martin & Co., New York City
42. Eastman Kodak Co., Rochester, N. Y.
43. Esonomic Materials Co., Chicago, Ill.
45. Petron Chem. Co., Brooklyn, N. Y.
46. Petron Chem. Co., Brooklyn, N. Y.
47. Fezandié and Sperilé, Inc., New York City
48. Fezandié and Sperilé, Inc., New York City
49. Fezandié and Sperilé, Inc., New York City
40. Fetron Chem. Co., Brooklyn, N. Y.
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59. Fries Bros., New York City
60. Fritzchie Bros., New York City
61. Geigy Co. Inc., New York City
62. General Atlas Carbon Co., New York City
63. General Chemical Co., New York City
64. General Dyestuffs Corp., New York City
65. General Electric Co., Schenectady, N. Y.
67. General Naval Stores Co., New York City
68. General Plastics Inc., No. Tonawands, N. Y.
68. General Plastics Inc., No. Tonawands, N. Y.
            of, General Naval Stores Co., New York City
68. General Plastics Inc., No. Tonawanda, N. Y.
69. Givaudan Delawanna, Inc., New York City
70. Glyco Products Co., Inc., New York City
71. Goldschmidt Corp., New York City
72. Goodyear Tire & Rubber Co., Akron, O.
73. Grasselli Chem. Co., Cleveland, O.
            73. Gressell Chem. Co., Clevenand, O.
74. Greef, R. W. & Co., Inc., New York City
75. Hall, C. P. & Co., Akron, O.
76. Halowax Corp., New York City
77. Harshaw Chem. Co., Cleveland, O.
78. Heine & Co., New York City
         78. Henne & Co., New York City
79. Hercules Powder Co., Wilmington, Del.
80. Hooker Electro-Chem. Co., New York City
81. Hopkins, J. L. & Co., New York City
82. Industrial Chem. Sales Co., New York City
83. Innis, Speiden & Co., New York City
84. International Pulp Corp., New York City
85. Johns-Manville Corp., New York City
86. Jungmann & Co., New York City
87. Kay-Fries Chomicals Inc. New York City
         80. Jugmann & Co., New York City
81. Kay-Fries Chemicals, Inc., New York City
82. Kessler Chem. Corp., New York City
83. Koppers Products Co., Pittsburgh, Pa.
90. Krobs Pigment & Color Corp., Newark, N. J.
91. Lehn & Fink Corp., New York City
92. Levin John D. Ly., New Hore, P. J.
            92. Lewis, John D., Inc., Providence, R. I
          93. Liquid Carbonic Corp., Chicago, Ill.
94. Lucidol Corp., Buffalo, N. Y.
         94. Lucidoi Corp., Burnad, N. Y.
95. Magnus, Mabee & Reynard, Inc., New York City
96. Mallinckrodt Chem. Works, St. Louis, Mo.
97. Martin, Dennis Co., Newark, N. J.
98. Mathieson Alkali Co., New York City
99. McCormick & Co., Baltimore, Md.
   199. McCormick & Co., Baitimore, Md.
100. Merck & Co. Inc., Now York City
101. Monsanto Chem. Works, St. Louis, Mo.
102. Moore-Munger, New York City
103. Mutual Chem. Co. of Amer., Newark, N. J.
104. National Aluminate Corp., Chicago, Ill.
105. National Aniline & Chem. Co., Buffalo, N. Y.
106. National Oil Products Co., Harrison, N. J.
107. National Regin Oil & Size Co. New York Cit.
107. National Rosin Oil & Size Co., New York City
108. Naugatuck Chem. Co., New York City
109. Neville Co., Pittsburgh, Pa.
110. New Jersey Zinc Sales Co., New York City
111. Nulomoline Co., New York City
112. Nuodex Products, Inc., Newark, N. J.
113. Onyx Oil & Chem. Co., Passaic, N. J.
114. Papermakers' Chem. Corp., Wilmington, Del.
115. Paramet Chem. Corp., Long Island City, N. Y.
116. Penick, S. B. & Co., New York City
117. Penn. Alcohol Corp., Phila., Pa.
119. Pfalts & Bauer, Inc., New York City
120. Phila. Quartz Co., Phila., Pa.
121. Plymouth Organic Labs., New York City
123. Pylam Products Co., New York City
123. Pauh, Robert Inc., Newark, N. J.
      107. National Rosin Oil & Size Co., New York City
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124. Reilly Tar & Chem. Corp., Indianapolis, Ind.
 125. Resinous Prod. & Chem. Co., Philadelphia, Pa.
  126. Resinox Corp., New York City
 127. Revertex Corp., New York City
  128. Robeson Process Co., New York City
  129. Rohm-Hass Chem. Co., Philadelphia, Pa.
 130. Royce Chem. Co., Carlton Hill, N. J.
131. Rubber Service Labs. Co., Akron, O.

132. Russia Cement Co., Gloucester, Mass.
133. Salomon, L. A. & Bro., New York City
134. Sandoz Chem. Works, New York City

  135. Scholler Bros., Inc., Philadelphia, Pa.
 136. Schliemann Co., Inc., New York City137. Scott, Bader & Co., London, England138. Seeley & Co., New York City
138. Seeley & Co., New York City
139. Sharples Solvents Corp., Philadelphia, Pa.
140. Shawinigan, Ltd., New York City
141. Sherwood Petroleum Co., Brooklyn, N. Y.
142. Silver, Geo., Import Co., New York City
143. Sonneborn, L. Sons, New York City
144. Southwark Mfg. Co., Camden, N. J.
145. Spencer-Kellogg Co., New York City
146. Stamford Rubber Supply Co., Stamford, Conn.
147. Stanco, Inc., New York City
148. Standard Oil Co. of Calif., San Francisco, Cal.
149. Standard Oil Co. of New Jersey, New York City
150. Stauffer Chem. Co., New York City
151. Stein-Hall & Co., Inc., New York City
152. Sun Oil Co., Philadelphia, Pa.
  152. Sun Oil Co., Philadelphia, Pa.
 153. Swann Chem. Corp., Birmingham, Ala.
154. Synfleur Scientific Labs., Monticello, N. Y.
 155. Texas Mining & Smelting Co., Laredo, Texas
156. Texas Mining & Smelting Co., Laredo, Texas
156. Thomas, Arthur H., Co., Philadelphia, Pa.
157. Titanium Pigments Co., New York City
158. Uhlich, Paul Co., New York City
159. United Color & Pigment Co., Inc., Newark, N. J.
 160. United States Gypsum Co., Chicago, Ill.
161. United States Industrial Chem. Co., Inc., New York City
161. United States Industrial Chem. Co., Inc., New York City 162. Van-Ameringen Haebler, Inc., New York City 163. Vanderbilt, R. T. Co., Inc., New York City 164. Varcum Chem. Corp., Niagara Falls, N. Y. 165. Verley, Albert & Co., Chicago, Ill. 166. Victor Chem. Works, Chicago, Ill. 167. Virginia Smelting Co., W. Norfolk, Va. 168. Vultex Corp. of America, Cambridge, Mass. 169. Wallestria Co. Lea Naw, Verk City.
 169. Wallerstein Co., Inc., New York City
170. Welch, Holme & Clark Co., Inc., New York City
171. Whittaker, Clark & Daniels, Inc., New York City
  172. Will & Baumer Candle Co., New York City
  173. Wishnick-Tumpeer, Inc., New York City
 174. Woburn Degreasing Co. of N. J., Harrison, N. J. 175. Wolf, Jacques & Co., Passaic, N. J. 176. Amer. Chemical Paint Co., Rochester, N. Y.
  177. Baker & Co., Inc., Newark, N. J.
 178. Chemical & Pigment Co., Baltimore, Md.
179. Heyden Chem. Works, New York, N. Y.
180. Kali Mfg. Co., Philadelphia, Pa.
181. Niacet Chem. Corp., Niagara Falls, N. Y.
182. Proctor & Gamble, Cincinnati, Ohio.
 183. Pure Calcium Products Co., Gainesville, O.184. Van Schaack Bros. Chem. Co., Chicago, Ill.185. Wyodak Chem. Co., Cleveland, O.
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WHERE TO BUY CHEMICALS

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Accelerators, Vulcanization
   Rubber Service Labs., Inc., Akron, O.
 Acetamide
   Amer. Chemical Products Co., Rochester, N. Y.
 Acetic Acid
   The Cleveland-Cliffs Iron Co., Cleveland, Ohio
 Acetic Anhydride
   American-British Chemical Supplies, Inc., New York, N. Y.
 Acetone
   W. S. Gray Co., New York, N. Y.
 Acetphenetidin
   Merck & Co., Inc., Rahway, N. J.
 Acetyl Balicylic Acid
   Monsanto Chemical Co., St. Louis, Mo.
 Acids, Fatty
Arthur C. Trask Co., Chicago, Ill.
 Acriflavine .
   Abbott Laboratories, North Chicago, Ill.
Agar
   American Agar Co., Inc., San Diego, Calif.
Albumen
   Stein, Hall & Co., Inc., New York, N. Y.
Alcohol, Denatured
  Rogers & McClellan, Boston, Mass.
L. R. Van Allen & Co., Chicago, Ill.
Alcohol, Pure
  U. S. Industrial Alcohol Co., New York, N. Y.
Alkalies
  Columbia Alkali Corp., New York, N. Y.
Alkaloids
  Merck & Co., Inc., Rahway, N. J.
Alkanet
  J. L. Hopkins & Co., New York, N. Y.
Almond Oil
  Magnus, Mabee & Reynard, Inc., New York, N. Y.
  Peck & Velsor, New York, N. Y.
Alpha Naphthol
Hord Color Products, Sandusky, O.
Alumina
  Aluminum Co. of America, Pittsburgh, Pa.
  Aluminum Co. of America, Pittsburgh, Pa.
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Abatio Acid

Hercules Powder Co., New York, N. Y.

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Aluminum Hydrate
    Ceramic Color & Chem. Mfg. Co., New Brighton, Pa.
  Alums
    The Grasselli Chemical Co., Cleveland, O.
  Aluminum Acetate
   Niacet Chemicals Corp., Niagara Falls, N. Y.
 Aluminum Bronze Powder
   U. S. Bronze Powder Works, Inc., New York, N. Y.
 Aluminum Chloride (Solution, Crystals and Anhydrous)
   The Calco Chemical Co., Bound Brook, N. J.
 Aluminum Stearate
   Franks Chemical Products Co., Inc., Brooklyn, N. Y.
   Glyco Products Co., Inc., New York, N. Y.
 Ammonia
   Nat'l Ammonia Co., Inc., Philadelphia, Pa.
 Ammonium Bifluoride
   The Harshaw Chemical Co., Cleveland, O.
 Ammonium Carbonate
   Wishnick-Tumpeer, Inc., New York, N. Y.
 Ammonium Chloride
   Pennsylvania Salt Mfg. Co., Inc., Philadelphia, Pa.
 Ammonium Linoleate
   Glyco Products Co., Inc., New York, N. Y.
 Ammonium Nitrate
   Garrigues, Stewart & Davies, Inc., New York, N. Y.
 Ammonium Oleate
  Glyco Products Co., Inc., New York, N. Y.
 Ammonium Persulphate
  Buffalo Electro Chemical Co., Inc., Buffalo, N. Y.
Ammonium Phosphate
  Swann Chemical Co., New York, N. Y.
Ammonium Sulphate
  H. J. Baker & Bro., New York, N. Y.
Ammonium Stearate
  Glyco Products Co., Inc., New York, N. Y.
Amyl Acetate
  Chemical Solvents, Inc., New York, N. Y.
Aniline Dyes
  Experimenter's Supply Co., New York, N. Y.
Aniline Oil
  Dow Chemical Co., Midland, Michigan
Antimony
  C. Tennant & Sons Co. of N. Y., New York, N. Y.
Antimony Chloride
  Seldner & Enequist, Inc., Brooklyn, N. Y.
Antimony Oxide
  O. Hommel Co., Pittsburgh, Pa.
Antimony Sulphide
Foote Mineral Co., Philadelphia, Pa.
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Anti-Oxidants

Givaudan-Delawanna, Inc., New York, N. Y.

Arsenio

Amer. Smelting & Refining Co., New York, N. Y.

Powhatan Mining Corp., Woodlawn, Baltimore, Md.

Asphalt

The Barber Asphalt Co., Philadelphia, Pa.

Asphaltum

Allied Asphalt & Mineral Corp., New York, N. Y.

Balsams

James B. Horner, Inc., New York, N. Y.

Barium Carbonate

Barium Reduction Corp., Charleston, W. Va.

Barium Nitrate C. W. Campbell Co., Inc., New York, N. Y.

Barium Peroxide

Barium Reduction Corp., Charleston, W. Va.

Barium Sulphate

C. P. De Lore Co., St. Louis, Mo.

Barium Sulphide

Chicago Copper & Chemical Co., Blue Island, Ill.

Barytes

Bradley & Baker, New York, N. Y. Nat'l Pigments & Chemical Co., St. Louis, Mo.

Basic Colors

Amer. Amline Products, Inc., New York, N. Y.

Bayberry Wax

The W. H. Bowdlear Co., Syracuse, N. Y.

A. C. Drury & Co., Inc., Chicago, Ill. Theodor Leonhard Wax Co., Inc., Haledon, Paterson, N. J.

Bentonite

Amer. Colloid Co., Chicago, Ill.

Silica Products Co., Kansas City, Mo. The Wyodak Chemical Co., Cleveland, Ohio

Benzaldchyde

Heyden Chem. Corp., New York, N. Y.

Bensidine

General Aniline Works, Inc., New York, N. Y.

Amer. Mineral Spirits Co., New York, N. Y.

Bensocaine

Abbott Laboratories, No. Chicago, Ill.

Bensoic Acid

Carus Chemical Co., Inc., La Salle, Ill.

Bensol

The Barrett Co., New York, N. Y.

Bensoul Peroxide

Lucidol Corp., Buffalo, N. Y.

Bensyl Cellulose

Advance Solvents & Chem. Corp., New York, N. Y.

Bergamot Oil

Orbis Products Corp., New York, N. Y.

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Beryllium
 Belmont Smelting & Refining Wks., Inc., Brooklyn, N. Y.
Beryllium and Its Salts
 Beryllium Corp. of America, New York, N. Y.
Beta Naphthol
 The Calco Chemical Co., Bound Brook, N. J.
 Cerro de Pasco Copper Corp., New York, N. Y.
Bismuth Subnitrate
 The New York Quinine & Chemical Wks., Inc., Brooklyn, N. Y.
 Adolph Hurst & Co., Inc., New York, N. Y.
Bleaching Powder
 Electro Bleaching Gas Co., New York, N. Y.
Blood Albumen
 Morningstar, Nicol, Inc., New York, N. Y.
Bone Ash
 Denver Fire Clay Co., Denver, Colorado
Bone Black
 Siemon Colors, Inc., Newark, N. J.
Bone Glue
 Darling & Co., Chicago, Ill.
Bone Oil
  Texas Chemical Co., Houston, Texas
 American Potash & Chem. Corp., New York, N. Y.
Bordeaux Mixture
 Mechling Bros. Chem. Co., Camden, N. J.
Borse Acid
 Borax Union, Inc., San Francisco, Calif.
Botanical Products
 S. B. Penick & Co., New York, N. Y.
 J. Q. Dickinson & Co., Malden, W. Va.
Bromo-Fluorescein
 Glyco Products Co., Inc., New York, N. Y.
Bronze Powder
 B. K. Drakenfeld & Co., New York, N. Y.
Burgundy Pitch
 Geo. H. Lincks, New York, N. Y.
Butyl Acetate
 Commercial Solvents Corp., New York, N. Y.
 Publicker, Inc., Philadelphia, Pa.
Butyl Aldehyde
  Commercial Solvents Corp., Terre Haute, Ind.
Butyl Alcohol (Normal)
 Publicker, Inc., Philadelphia, Pa.
Butyl Propionate
 C. P. Chemical Solvents, Inc., New York, N. Y.
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The Northwestern Chemical Co., Wauwatosa, Wisconsin

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Butul Stearate
  Kessler Chem. Corp., New York, N. Y.
Cadmiu#
  U. S. Smelting, Refining & Mining Co., New York, N. Y.
Cajuput Oil
  D. W. Hutchinson & Co., New York, N. Y.
Calcium Arsenate
  Bowker Chemical Corp., New York, N. Y. Chipman Chemical Co., Inc., Bound Brook, N. J.
Calcium Carbonate
  Limestone Products Corp. of Amer., Newton, N. J.
Calcium Carbonate (Precipitated)
  Merck & Co., Inc., Rahway, N. J.
Calcium Chloride
  Michigan Alkali Co., New York, N. Y.
  Saginaw Salt Products Co., Saginaw, Mich.
Calcium Chloride (Anhydrous)
Fales Chemical Co., Inc., Cornwall Landing, N. Y.
Calcium Phosphate
  Provident Chemical Wks., St. Louis, Mo.
Calcium Sulphide (Luminous)
  Amer. Luminous Products Co., Huntington Park, Calif.
Calcium Stearate
  The Synthetic Products Co., Cleveland, Ohio
Camphor
  E. J. Barry, New York, N. Y.
Camphor Oil
  Magnus, Mabee & Reynard, Inc., New York, N. Y.
Candelilla Wax
  Innis, Speiden & Co., Inc., New York, N. Y.
Caramel Color
  Alex Fries & Bro., Cincinnati, Ohio
Caraway Oil
  Geo. Lueders & Co., New York, N. Y.
Carbolic Oil
  Reilly Tar & Chemical Corp., New York, N. Y.
Carbon, Activated
  The Jennison-Wright Co., Toledo, Ohio
Carbon Bisulphide
  J. T. Baker Chemical Co., Phillipsburg, N. J.
Carbon Black
  United Carbon Co., Charleston, W. Va.
Binney & Smith, New York, N. Y.
Carbon, Decolorizing
  Darco Sales Corp., New York, N. Y.
Carbon Tetrachloride
  Niagara Smelting Corp., Niagara Falls, N. Y.
Cardamom Seed
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Casein
The Casein Mfg. Co. of America, Inc., New York, N. Y.

Newmann-Buslee & Wolfe, Inc., Chicago, Ill.

Frank B. Ross Co., Inc., New York, N. Y.

Carnauba Wax

Castile Soap Conti Products Corp., New York, N. Y. Castor Oil The Baker Castor Oil Co., New York, N. Y. Castor Oil, Sulphonated Jacques Wolf & Co., Passaic, N. J. Celluloid Celluloid Corp., New York, N. Y. Celluloid Scrap Moses Serinsky Co., Indianapolis, Ind. Cellulose Acetate Celanese Corp. of America, New York, N. Y. Cellulose Nitrate Merrimac Chemical Co., Everett, Mass. Ceresin Wax Sherwood Petroleum Co., Inc., Brooklyn, N. Y. Cetyl Alcohol Hummel Chemical Co., Inc., 90 West St., New York, N. Y. Chalk, Precipitated Charles B. Chrystal Co., Inc., New York, N. Y. Chas. L. Read & Co., Inc., New York, N. Y. Western Charcoal Co., Chicago, Ill. China Clay Taintor Trading Co., New York, N. Y. China Wood Oil Balfour, Guthrie & Co., Ltd., New York, N. Y. Chloramine Abbott Laboratories, No. Chicago, Ill. Chlorine (Liquid) Electro Bleaching Gas Co., 9 E. 41st St., New York, N. Y. The Dow Chemical Co., Midland, Michigan Chlorophyll Amer. Chlorophyll, Inc., New York, N. Y. Pylam Products Co., New York, N. Y. Cholesterin Digestive Ferments Co., Detroit, Michigan Merck & Co., Inc., Rahway, N. J. Chrome Green Kentucky Color & Chem. Co., Louisville, Ky. Chrome Yellow Ansbacher-Siegle Corp., Rosebank, N. Y. Chromic Acid Mutual Chemical Co. of America, New York, N. Y. Chromium Oxide O. Hommel Co., Inc., Pittsburgh, Pa. Citral Givaudan-Delawanna, Inc., New York, N. Y. Citric Acid

Chas. Pfizer & Co., Inc., New York, N. Y.

H. C. Ryland, Inc., New York, N. Y.

Citronella Oil

Coumarin

Coumarone Resin

Maywood Chem. Works, Maywood, N. J.

Barrett Co., New York, N. Y. Neville Co., Pittsburgh, Pa.

Clay
Kentucky Clay Mining Co., Mayfield, Ky.
Olive Branch Minerals Co., Cairo, Ill. Coal Ta Crowley Tar Products Co., New York, N. Y. Coal Tar Colors H. Kohnstamm & Co., New York, N. Y. Cobalt Acetate Fred L. Brooke Co., Chicago, Ill. Cobalt Driers McGean Chemical Co., Cleveland, Ohio Cobalt Linoleate The McGean Chemical Co., Cleveland, Ohio Cocoa Butter Alpha Lux Co., Inc., New York, N. Y. Thomas J. Shields Co., New York, N. Y. Coconut Butter Procter & Gamble Co., Cincinnati, Ohio Coconut Oil Franklin Baker Co., Hoboken, N. J. Coconut Oil Fatty Acid Aeme Oil Corp., Chicago, Ill. Cod Liver Oil H. H. Rosenthal & Co., Inc., New York, N. Y. Charles Cooper & Co., New York, N. Y. Colors, Dry Holland Aniline Dye Co., Holland, Mich. Colors, Oil Soluble Commonwealth Color & Chem. Co., Brooklyn, N. Y. Copper Carbonate Chas. Copper & Co., New York, N. Y. Jungmann & Co., Inc., New York, N. Y. Copper Cyanide Charles Hardy, Inc., New York, N. Y. Copper Oxides The O. Hommel Co., Inc., 209 Fourth Ave., Pittsburgh, Pa. Copper Sulphate Barada & Page, Inc., Kansas City, Mo. Corn Oil American Maize Products Co., New York, N. Y. Corn Sugar Staley Sales Corp., Decatur, Ill. Corn Syrup Clinton Co., Clinton, Ia. Corn Products Refining Co., New York, N. Y. Cottonseed Oil (Crude) Battleboro Oil Co., Battleboro, N. C. Welch, Holme & Clark Co., New York, N. Y.

Cream of Tartar
The Harshaw Chemical Co., Cleveland, Ohio

Creosote
Koppers Products Co., Pittsburgh, Pa.

resole

Coopers Creek Chem. Co., W. Conshohocken, Pa. Reilly Tar & Chemical Corp., New York, N. Y.

Cresylic Acid
The Barrett Co., New York, N. Y.

Cryolite
Vitro Mfg. Co., Pittsburgh, Pa.

Cyclohexanol

E. I. Du Pont de Nemours Co., Wilmington, Del.

Geo. H. Lincks, New York, N. Y.

Amer. Lanolin Corp., Lawrence, Mass.

Derris Extract
Seacoast Laboratories, New York, N. Y.

W. Benkert & Co., Inc., New York, N. Y.

Dextrins
Morningstar, Nicol, Inc., New York, N. Y.

Dustase
Takamine Laboratory, Inc., Clifton, N. J.

Diatomaceous Earth
Dicalite Co., New York, N. Y.

Dibutylphthalate
The Kessler Chemical Corp., New York, N. Y.

Dichlorbenzol
Hooker Electro Chemical Co., New York, N. Y.

Dicthyleneglycol Carbide & Carbon Chemicals Corp., New York, N. Y.

Diethylphthalate Van Dyk & Co., Inc., Jersey City, N. J.

Digiycol Oleate
Glyco Products Co., Inc., New York, N. Y.

Diglycol Laurate
Glyco Products Co., Inc., New York, N. Y.

Diglycol Stearate
Glyco Products Co., Inc., New York, N. Y.

Carbide & Carbon Chem. Corp., New York, N. Y.

Dipentene Hercules Powder Co., Wilmington, Del.

Diphenyl Swann Chemical Co., New York, N. Y.

Drop Black
Wilckes-Martin-Wilckes Co., New York, N. Y.

Dyestuffs
National Aniline & Chemical Co., Inc., New York, N. Y.

Egg, Dried *
W. P. Pray, New York, N. Y.

Egg Yolk Stein, Hall & Co., New York, N. Y.

Ephedrine
Abbott Laboratories, No. Chicago, Ill.

Epsom Salt
General Chemical Co., New York, N. Y.

Essential Oils
Compagnie Duval, New York, N. Y.

Compagnie Duvai, New York, N. 1.
Ester Gum

John D. Lewis, Inc., Providence, R. I. Paramet Chemical Corp., Long Island City, N. Y.

Carbide & Carbon Chemicals Corp., New York, N. Y.

Ethyl Acetate
Merrimac Chemical Co., Boston, Mass.

Ethyl Cellulose
Advance Solvents & Chem. Corp., New York, N. Y.

F. C. Bersworth Labs., Framingham, Mass.

Ethyl Lactate
American Cyanamid & Chemical Corp., New York, N. Y.

Ethylene Diamine F. C. Bersworth Labs., Framingham, Mass.

Ethylene Dichloride Dow Chemical Co., Midland, Mich.

Ethyleneglycol
Carbide & Carbon Chemicals Corp., New York, N. Y.

Eucalyptus Oil Chas. Fishbeck Co., New York, N. Y.

Feldspar Consolidated Feldspar Corp., Trenton, N. J.

Fillers C. K. Williams & Co., Easton, Pa.

Film Scrap Horn-Jefferys & Co., Burbank, Calif.

Fish Glue C. B. Hewitt & Bro., New York, N. Y.

Fish Oil Falk & Co., Pittsburgh, Pa.

Flaxseed
Bisbee Linseed Co., Philadelphia, P.

Fluorspar Hillside Fluor Spar Mines, Chicago, Ill.

Formic Acid Victor Chem. Works, Chicago, Ill.

Victor Chem. Works, Chicago, Ill. Formaldehyde

Heyden Chemical Corp., New York, N. Y.

Fuller's Earth
L. A. Salmon & Bro., New York, N. Y.
Sinclair Refining Co., Olmstead, Ill.

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Fusel Oil
  Empire Distilling Corp., New York, N. Y.
Gallio Acid
  Eastman Kodak Co., Rochester, N. Y.
Gamboge
  Frank B. Ross Co., New York, N. Y.
  Atlantic Gelatine Co., Woburn, Mass.
Geraniol
  Kay-Fries Chem., Inc., New York, N. Y.
Geranium Lake
  Interstate Color Co., Inc., New York, N. Y.
  R. F. Revson Co., New York, N. Y.
Geranium Oil
  Schimmel & Co., New York, N. Y.
Gilsonite
  George H. Lincks, New York, N. Y.
  Utah Gilsonite Co., St. Louis, Mo.
Ginseng
C. H. Lewis & Co., New York, N. Y.
Glandular Products
  The Wilson Laboratories, Chicago, Ill.
  Iowa Soda Products Co., Council Bluffs, Is.
  Cudahy Packing Co., Chicago, Ill.
Glycerin
  Colgate-Palmolive-Peet Co., Chicago, Ill.
Glyceryl Mono Stearate
  Glyco Products Co., Inc., New York, N. Y.
Glyceryl Phthalate
  Glyco Products Co., Inc., New York, N. Y.
Glyceryl Stearate
  Glyco Products Co., Inc., New York, N. Y.
  Glyco Products Co., Inc., New York, N. Y.
Glycol Phthalate
  Glyco Products Co., Inc., New York, N. Y.
Glycol Stearate
  Glyco Products Co., Inc., New York, N. Y.
Gold Chloride
  Mallinckrodt Chemical Works, St. Louis, Mo.
  Adolphe Hurst & Co., Inc., New York, N. Y.
Asbury Graphite Mills, Asbury Park, N. J.
Gum Arabic
  T. M. Duche & Sons, New York, N. Y.
Gum Bensoin
 Peek & Velsor, Inc., New York, N. Y.
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Gum Copal

Gum Damar

George H. Lincks, New York, N. Y.

Thurston & Braidich, New York, N. Y.

Gum Karaya
Frank-Vliet Co., Inc., New York, N. Y.
Gum, Locust Bean

Innis, Speiden Co., New York, N. Y.

Gum Manila Stroock & Wittenberg Corp., New York, N. Y.

Gum Tragacanth
E. Meer & Co., Inc., New York, N. Y.
J. L. Hopkins & Co., New York, N. Y.

Gypsum U. S. Phosphoric Prod. Corp., New York, N. Y.

Hemlock Bark
Tanners Supply Co., Grand Rapids, Mich.

Henna Leaves
S. B. Penick & Co., New York, N. Y.

Herbs

John Clarke & Co., New York, N. Y. Hexamethylenetetramine

Heyden Chemical Corp., New York, N. Y. Hydrochlorio Acid General Chemical Co., New York, N. Y.

Hydrogen Peroxide
The Warner Chemical Co., New York, N. Y.

Hydroquinone Eastman Kodak Co., Rochester, N. Y.

Ichthyol

Merck & Co., Rahway, N. J.

Indigo L. E. Ransom Co., New York, N. Y.

Belmont Smelting & Refining Works, Brooklyn, N. Y.

Invert Sugar Nulomoline Co., New York, N. Y.

Iodine New York Quinine & Chemical Wks., Inc., Brooklyn, N. Y.

Iridium
Baker & Co., Inc., Newark, N. J.

Baker & Co., Inc., Newark, N. J. Irish Moss

S. B. Penick & Co., New York, N. Y. Iron Ammonium Citrate

Schuykill Chem. Co., Philadelphia, Pa.

Iron Chloride

Chicago Copper & Chem. Co., Blue Island, Ill.

Iron Oxide
Binney & Smith Co., New York, N. Y.

Isopropyl Acetate
A. K. Hamilton, New York, N. Y.

Isopropyl Alcohol
Carbide & Carbon Chemicals Corp., New York, N. Y.

Insect Wax, Chinese Frank B. Ross Co., Inc., New York, N. Y.

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Ivory Black
     Binney & Smith Co., New York, N. Y.
  Japan Wax
     Smith & Nichols, Inc., New York, N. Y.
  Kerosens
     Colonial Beacon Oil Co., Everett, Mass.
  Kerosene, Deodorized
     Sherwood Petroleum Co., Brooklyn, N. Y.
 Laboratory Equipment
Central Scientific Co., Chicago, Ill.
Chemical Publ. Co. of N. Y., Inc., New York, N. Y.
Chicago Apparatus Co., Chicago, Ill.
Eimer & Amend, New York, N. Y.
Experimenter's Supply Co., New York, N. Y.
Fisher Scientific Co., Pittsburgh, Pa.
N. J. Laborators Kundy Co. Newark N. J.
    N. J. Laboratory Supply Co., Newark, N. J.
Scientific Glass Apparatus Co., Bloomfield, N. J.
 Lacquers
    Maas & Waldstein, Newark, N. J.
    Apex Chemical Co., Inc., New York, N. Y.
 Lamp Black
    Binney & Smith Co., New York, N. Y.
L. Martin Co., New York, N. Y.
 Lanolin
   American Lanolin Corp., Lawrence, Mass.
Merck & Co., Inc., Rahway, N. J.
Pfaltz & Bauer, New York, N. Y.
 Lard Oil
   Enterprise Animal Oil Co., Philadelphia, Pa.
 Lauryl Alcohol and Sulphonate
   E. I. Du Pont de Nemours & Co., Wilmington, Del.
 Lavender Oil
    Van Ameringen-Haebler, Inc., New York, N. Y.
Lead Acetate
   National Lead Co., New York, N. Y.
Lead Arsenate
   Barada & Page, Inc., Kansas City, Mo. General Chemical Co., New York, N. Y.
Lead and Its Oxides
   The Eagle-Picher Sales Co, Cincinnati, Ohio
Lecithin
   American Lecithin Corp., New York, N. Y.
Lemon Juice, Concentrated
   Mutual Citrus Products Co., Anaheim, Calif.
   D. W. Hutchinson & Co., Inc., New York, N. Y.
Licorice
  MacAndrews & Forbes Co., New York, N. Y.
Lime
  J. E. Baker Co., York, Pa.
Chazy Marble Lime Co., Inc., Chazy, N. Y.
Limestone
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F. E. Schundler & Co., Joliet, Ill.

Linoleic Acid Glyco Products Co., Inc., New York, N. Y.

Linseed Oil Bisbee Linseed Co., Philadelphia, Pa.

Litharge
The Eagle-Picher Lead Co., Cincinnati, Ohio

Lithopone
Krebs Pigment & Color Corp., Newark, N. J.
Marshall Dill Co., San Francisco, Calif.

Locust Bean Powder
T. M. Duche & Sons, New York, N. Y.

Logwood Extract
American Dyewood Co., New York, N. Y.

Lycopodium

McKesson & Robbins, Inc., New York, N. Y.

Magnesia Philip Carey Co., Lockland, O.

Magnesite
General Magnesite & Magnesia Co., Philadelphia, Pa.

Magnesium Carbonate Merck & Co., Inc., Rahway, N. J.

Magnesium Chloride Wishnick-Tumpeer, Inc., New York, N. Y.

Magnesium Powder
Belmont Smelting & Refining Wks., Inc., Brooklyn, N. Y.

Maleic Acid Nat'l Aniline & Chem. Wks., New York, N. Y.

Manganese
Ajax Metal Co., Philadelphia, Pa.

Marble Dust Hammil & Gillespie, Inc., New York, N. Y.

Manganese Dioxide
B. F. Drakenfeld & Co., Inc., New York, N. Y.

Menhaden Oil Robert Badcock & Co., New York, N. Y.

Menthol
Chas. L. Huisking & Co., Inc., New York, N. Y.

Mercury
Chas. L. Huisking & Co., Inc., New York, N. Y.
George Uhe Co., New York, N. Y.

Methanol
Wm. S. Gray & Co., New York, N. Y.

Methyl Acetate
Carbide & Carbon Chem. Corp., New York, N. Y.

Methyl Acctone
Delta Chem. & Iron Co., Wells, Mich.

Methyl Anthranilate
Florasynth Laboratories, New York, N. Y.

Methyl p-Hydroxybensoate
Heyden Chemical Corp., New York, N. Y.

Methyl Salicylate
Dow Chemical Co., Midland, Michigan

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Southern Mica Co., Franklin, N. C.
 Milk Sugar
   Mallinckrodt Chemical Wks., St. Louis, Mo.
Mineral Rubber
  Barber Asphalt Co., Philadelphia, Pa.
Mineral Spirits
 . Amer. Mineral Spirit Co., New York, N. Y.
Montan Wax
  Strahl & Pitsch, New York, N. Y.
Naphtha
  Deep Rock Oil Corp., Chicago, Ill.
Naphthalene
  The Barrett Co., New York, N. Y.
Naphthenio Acid
  Glyco Products Co., Inc., New York, N. Y.
Neatsfoot Oil
  National Oil Products Co., Harrison, N. J.
Nickel Chloride
  Chas. Cooper & Co., New York, N. Y.
Nickel Sulphate
  The Harshaw Chemical Co., Cleveland, O.
Nicotine
  Tobacco By-Products & Chemical Corp., Louisville, Ky.
Nicotine Sulphate
  Lattimer-Goodwin Chemical Co., Grand Junction, Colo.
Nitre Cake
  Trojan Powder Co., Allentown, Pa.
Nitric Acid
  Monsanto Chemical Co., St. Louis, Mo.
Nitrobensol
  Calco Cher. Co., Bound Brook, N. J.
Nitrocellulose
  E. I. Du Pont de Nemours & Co., Inc., Parlin, N. J.
 Smith Chemical & Color Co., Brooklyn, N. Y.
Oil, Citronella
 D. W. Hutchinson & Co., Inc., New York, N. Y.
Oil, Mineral
 Standard Oil Co. of California, San Francisco, Calif.
Oil, Olive
 Leghorn Trading Co., Inc., New York, N. Y.
Oiticica Oil
 L. N. Jackson & Co., New York, N. Y.
Olein
 Century Stearic Acid Wks., New York, N. Y.
Oleoresins
 Seeley & Co., New York, N. Y.
Olive Oil, Sulphonated
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Royce Chem. Co., Carlton Hill, N. J.

Dodge & Olcott Co., New York, N. Y.

Orange Oil

Phosphoric Acid

Victor Chemical Works, Chicago, Ill.

Ortho Dichlorbenzene Hooker Electrochemical Co., New York, N. Y. Oxalic Acid Mutual Chemical Co. of America, New York, N. Y. Wilson Labs., Chicago, Ill. Oxygen Cheney Chemical Co., Cleveland, O. Oxyguinoline Sulphate Benzol Products Co., Newark, N. J. Ozokerite Wax Strohmeyer & Arpe Co., New York, N. Y. Palm Kernel Oil Franklin Baker Co., Hoboken, N. J. Palm Oil Wishnick-Tumpeer, Inc., New York, N. Y. Paraffin Oils S. Schwabacher & Co., Inc., New York, N. Y. Paraffin Wax Oil States Petroleum Co., New York, N. Y. Paraldehyde Heyden Chem. Corp., New York, N. Y. Para Aminophenol Verona Chem Co., Newark, N. J. Para-Phenylenediamine Amido Products Co., New York, N. Y. Paris White Southwark Mfg. Co., Camden, N. J. Peanut Oil Elbert & Co., New York, N. Y. Pearl Essence Mearl Corp., New York, N. Y. Pectin Calif. Fruit Growers' Exchange, Ontario, Calif. Peppermint Oil Magnus, Mabee & Reynard, Inc., New York, N. Y. The Sparhawk Co., Sparkhill, N. Y. Perilla Oil S. L. Jones & Co., San Francisco, Calif. Pennsylvania Refining Co., Butler, Pa. Petroleum Jelly L. Sonneborn Sons, Inc., New York, N. Y. Petrolcum Spirits Sun Oil Co., Philadelphia, Pa. American-British Chemical Supplies, Inc., New York, N. Y. Phenol-Formaldehyde Resins Durite Plastics, Philadelphia, Pa.

Phosphorus

International Selling Corp., New York, N. Y.

Phthalic Anhydride

Monsanto Chem. Co., St. Louis, Mo.

General Naval Stores Co., Inc., New York, N. Y.

Pine Tar Southern Pine Chem. Co., Jacksonville, Fla.

Robert Rauh, Inc., Newark, N. J.

Plaster of Paris

Whittaker, Clark & Daniels, Inc., New York, N. Y.

Potash, Caustio

Niagara Alkali Co., New York, N. Y.

Potassium Carbonate

Joseph Turner & Co., New York, N. Y.

Potassium Chlorate Joseph Turner & Co., New York, N. Y.

Potassium Hydroxide

Merck & Co., Inc., Rahway, N. J.

New York Quinine & Chemical Wks., Inc., Brooklyn, N. Y.

Glyco Products Co., Inc., New York, N. Y. Carl F. Miller & Co., Scattle, Washington

Potassium Permanganate

Carus Chemical Co., Inc., La Salle, Ill.

Potassium Silicate

Philadelphia Quartz Co., Philadelphia, Pa.

Prussian Blue Kentucky Color & Chem. Co., Louisville, Ky.

Charles B. Crystal Co., New York, N. Y.

Psyllium Seeds Laxseed Co., New York, N. Y.

Pyrethrum Extract

McLaughlin, Gormley, King & Co., Minneapolis, Minn.

Pyrethrum

S. B. Penick & Co., New York, N. Y.

Pyrogallic Acid

Zinsser & Co., Inc., Hastings on Hudson, N. Y.

Pyroxylin Solutions

Egyptian Lacquer, Kearney, N. J.

Quince Seed

J. L. Hopkins & Co., New York, N. Y.

Quinine Bisulphate

R. W. Greef & Co., Inc., New York, N. Y.

Rapeseed Oil

Balfour, Guthrie & Co., Ltd., New York, N. Y.

Century Stearic Acid Candle Wks., New York, N. Y.

Resins, Synthetic Beck, Koller & Co., Inc., Detroit, Mich. Marshall Dill, San Francisco, Calif.

Penn. Coal Products Co., Petrolia, Pa.

Rhodium Baker & Co., Inc., Newark, N. J.

Rochelle Salts

Chas. Pfizer & Co., Inc., New York, N. Y.

Rose Water

Geo. Lueders & Co., New York, N. Y.

Rosin General Naval Stores Co., Inc., New York, N. Y.

National Rosin Oil & Size Co., New York, N. Y.

Rotenone

Thorocide, Inc., St. Louis, Mo.

Rubber

Earle Bros., New York, N. Y.

Rubber Latex

Littlejohn & Co., Inc., New York, N. Y.

Saccharine

Heyden Chemical Corp., New York, N. Y.

Salicylic Acid

The Dow Chemical Co., Midland, Mich.

Sal Soda Church & Dwight Co., Inc., New York, N. Y.

Morton Salt Co., Chicago, Ill.

Amer. Cyanamid & Chem. Corp., New York, N. Y.

Croton Chem. Corp., Brooklyn, N. Y.

Saponin

Experimenters Supply Co., New York, N. Y. Jungmann & Co., New York, N. Y.

Selenium

Amer. Metal Co., New York, N. Y.

Wm. Zinsser & Co., New York, N. Y.

Adolphe Hurst & Co., New York, N. Y.

Fezandie & Sperrie, Inc., New York, N. Y.

Barnsdall Tripoli Corp., Seneca, Mo.

Silver

Handy & Harman, New York, N. Y.

Bilver Cyanide Chas. Cooper & Co., New York, N. Y.

Silver Nitrate

Zastman Kodak Co., Rochester, N. Y.

Soda Ash

Diamond Alkali Co., Pittsburgh, Pa.

Soda, Caustic

Mathieson Alkali Works, Inc., New York, N. Y.

Soda, Sal

Consolidated Chema Sales Corp., Newark, N. J.

Sodium Aluminate

National Aluminate Corp., Chicago, Ill.

Sodium Arsenite

Harrison Mfg. Co., Rahway, N. J.

Sodium Benzoate

Hooker Electrochemical Co., New York, N. Y.

Sodium Bicarbonate

Church & Dwight Co., Inc., New York, N. Y.

Sodium Bichromate Prior Chem. Corp., New York, N. Y.

Sodium Bisulphite

The Grasselli Chemical Co., Cleveland, Ohio

Sodium Carbonate

Solvay Sales Corporation, New York, N. Y.

Sodium Cholcate

Difco Laboratories, Inc., Detroit, Mich.

Sodium Cyanide

E. I. Du Pont de Nemours & Co., Inc., Wilmington, Del.

Sodium Fluoride

American Cyanamid & Chemical Corp., New York, N. Y.

Sodium Hydrosulphite

Royce Chemical Co., Carlton Hill, N. J.

Sodium Hydroxide

Merck & Co., Inc., Rahway, N. J.

Sodium Hypochlorite

Delta Chemical Mfg. Co., Baltimore, Md.

Mathieson Alkali Wks., Inc., New York, N. Y.

Sodium Hypochlorite Liquid

Riverside Chemical Co., No. Tonawanda, N. Y.

Sodium Hyposulphite

The Grasselli Chemical Co., Cleveland, Ohio

Sodium Metaphosphate

Buromin Co., Pittsburgh, Pa.

Sodium Metasilicate

Philadelphia Quartz Co., Philadelphia, Pa.

Sodium Nitrate

Battelle & Renwick, New York, N. Y.

Sodium Nitrite

Solvay Sales Corp., New York, N. Y.

Sodium Perborate

E. I. Du Pont de Nemours & Co., Inc., Wilmington, Del.

Sodium Phosphate

Swann Chemical Co., New York, N. Y.

Sodium Resinate

Paper Makers Chem. Corp., Wilmington, Del.

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Sodium Silicate
Mechling Bros. Chemical Co., Camden, N. J.
Philadelphia Opartz Co., Philadelphia Pa

Philadelphia Quartz Co., Philadelphia, Pa. Standard Silicate Co., Pittsburgh, Pa.

Sodium Silico Fluoride
The Grasselli Co., Cleveland, Ohio

Sodium Sulphate General Chem. Co., New York, N. Y.

Sodium Stannate
Harshaw Chem. Co., Cleveland, Ohio

Sodium Sulphite
Mechling Bros. Chemical Co., Camden, N. J.

Sodium Tungstate
J. T. Baker Chem. Co., Phillipsburg, N. J.

Solvent Naphtha Barrett Co., New York, N. Y.

Sorbitol
Atlas Powder Co., Wilmington, Del.

Soybean Oil Spencer Kellogg & Sons Sales Corp., Buffalo, N. Y. Arthur C. Trask Co., Chicago, Ill.

Sperm Oil Cook Swan Co., Inc., New York, N. Y.

Spermaceti Strahl & Pitsch, New York, N. Y.

Squill
S. B. Penick & Co., New York, N. Y.

Starch Products Co., New York, N. Y.

Stearie Acid
Century Stearie Acid Candle Wks., New York, N. Y.

Stearin M. Werk Co., Cincinnati, Ohio

Stearine Pitch
A. Gross & Co., New York, N. Y.

Strontium Nitrate Grasselli Chem. Co., Cleveland, Ohio

Strychnine Chas. Pfizer & Co., New York, N. Y.

Sulphonated Castor Osl Burkard-Schier Chem. Co., Chattanooga, Tenn.

Sulphonated Olive Osl Jacques Wolf & Co., Passaic, N. J.

Sulphur Stauffer Chemical Co. of Texas, Freeport, Tex.

Sulphur Dioxide Virginia Smelting Co., Boston, Mass.

Sulphurio Acid
Merrimac Chemical Co., Everett Sta., Boston, Mass.

Talo Charles B. Crystal Co., Inc., New York, N. Y.

Welch, Holme & Clark Co., Inc., New York, N. Y. Tartaric Acid R. W. Greeff & Co., Inc., New York, N. Y. Tar Acid Oil Barrett Co., New York, N. Y. Tartar Emetic Apex Chem. Co., New York, N. Y. Tea Seed Oil Lundt & Co., New York, N. Y. D. W. Hutchinson & Co., New York, N. Y. Tetrachlorethane Dow Chemical Co., Midland, Mich. Tetrachlorethylene E. I. Du Pont de Nemours & Co., Wilmington, Del. Thallsum Sulphate Jungmann & Co., Inc., New York, N. Y. Thiocarbamilid Monsanto Chemical Co., St. Louis, Mo. Thiourca Jungmann & Co., New York, N. Y. Thymol Sherka Chemical Co., Inc., Bloomfield, N. J. Union Smelting & Refining Co., Inc., Newark, N. J. Tin Chloride Seldner & Enequist, Inc., Brooklyn, N. Y. Tin Oxide McGean Chemical Co., Cleveland, Ohio Tinctures Parke, Davis & Co., Detroit, Mich. Titanium Dioxide Marshall Dill, San Francisco, Calif. R. T. Vanderbilt Co., New York, N. Y. Toluol Jones & Laughlin Steel Corp., Pittsburgh, Pa. Triacetin Niacet Chemicals Corp., Niagara Falls, N. Y. Tricresyl Phosphate R. W. Greeff & Co., Inc., New York, N. Y. TriethanolamineExperimenter's Supply Co. (small lots), New York, N. Y. Carbide & Carbon Chem. Co. (large lots), New York, N. Y. Triethanolamine Oleate Glyco Products Co., Inc., New York, N. Y. Marshall Dill Co., San Francisco, Calif. Triethanolamine Stearate Glyco Products Co., Inc., New York, N. Y.

Carl F. Miller & Co., Seattle, Washington

Triphenylquanadine
E. I. Du Pont de Nemours & Co., Wilmington, Del.

Triphenylphosphate
Monsanto Chemical Co., St. Louis, Mo.

Tripoli
Tamms Silica Co., Chicago, Ill.

Tungsten
Fansteel Products Co., No. Chicago, Ill.

Turkey Red Oil
National Oil Products Co., Inc., Harrison, N. J.

Turpentine

Antwerp Naval Stores Co., Inc., Boston, Mass. General Naval Stores Co., New York, N. Y.

Turpentine Substitute
Anderson Prichard Oil Corp., Oklahoma City, Okla.

Turpentine (Venice)
National Rosin Oil & Size Co., New York, N. Y.

Turtle Oil

Edwin Seebach Co., New York, N. Y.

Ultramarine Blue

Standard Ultramarine Co., Huntington, W. Va. Umbers

Fezandie & Sperrle, Inc., New York, N. Y.

Uranium Nitrate
Harshaw Chemical Co., Cleveland, Ohio

Sherka Chemical Co., Inc., Bloomfield, N. J.

Vanilla Beans
Thurston & Braidich, New York, N. Y.

Vanillin Seelev &

Urea

Seeley & Co., Inc., New York, N. Y. Van Ameringen-Haebler, Inc., New York, N. Y.

Varnish Gums and Resins
Amer. Cyanamid & Chem. Corp., New York, N. Y.

Vat Colors
Amer. Aniline Products, Inc., New York, N. Y.

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L. E. Ransom Co., New York, N. Y.

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Hill Bros. Chem. Co., Los Angeles, Calif.

Vermilion Fezandié & Sperrlé, Inc., New York, N. Y.

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Niagara Chemicals Corp., Niagara Falls, N. Y.

Vinyl Chloride
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Wax, Synthetic Glyco Products Co., Inc., New York, N. Y.

Wetting Out Agents
Glyco Products Co., Inc., New York, N. Y.

Whiting
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National Lead Co., New York, N. Y.

Wood Flour
D. H. Litter Co., New York, N. Y.
Wood Flour, Inc., Manchester, N. H.

Yylol
The Barrett Co., New York, N. Y.

Yeast Standard Brands, Inc., New York, N. Y.

Zinc Hegeler Zinc Co., Danville, Ill.

Zinc Carbonate
Wishnick-Tumpeer, Inc., New York, N. Y.

Zinc Chloride
Wishnick-Tumpeer, Inc., New York, N. Y.

Zinc Chromate
E. M. & F. Waldo, Inc., Muirkirk, Md.

Zino Oxide Merck & Co., Inc., Rahway, N. J. N. J. Zinc Co., New York, N. Y.

Zinc Stearate
Merck & Co., Inc., Rahway, N. J.
Wishnick-Tumpeer, Inc., New York, N. Y.

Zinc Sulphate
W. R. Russell & Co., New York, N. Y.
Virginia Smelting Co., West Norfolk, Va.

Zirconium Oxide Foote Mineral Co., Philadelphia, Pa.

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       Estate, Bombay
   Ciba (India), Ltd., Post Box 479, Bombay
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W. A. Scholten's Chemical Works, Ltd., Groningen, Holland
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       27 Chesser St., Adelaide
414 Kent Ave., Sidney
19 Lower Tory St., Wellington, New Zealand
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    W. H. Goetz, Calle Sarandi 315, Buenos Aires
Cuba
   J. M. Sierra, Aquiar 73 Dpt. 710, Apartado 362, Havana
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